

Model-based investigation of nitrogen and oxygen cycles  
in the oxygen minimum zone of the eastern tropical South  
Pacific

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## Summary

Nitrogen is an essential nutrient for controlling marine primary production. Its cycling in the ocean is tightly associated with primary gain and loss processes, including nitrogen fixation, atmospheric deposition, water-column and benthic denitrification, and anaerobic ammonium oxidation (anammox). Oxygen minimum zones (OMZs) in the ocean are usually defined to be water columns with  $O_2$  concentration below  $20 \mu\text{mol kg}^{-1}$ . OMZs currently account for only about 8 % of the global ocean area, but could be responsible for as much as 30–50 % of the total fixed-N loss, mainly in form of  $N_2$ , into the atmosphere by denitrification or anammox. The eastern tropical South Pacific (ETSP), encompassing one of the permanent OMZs in the ocean, is responsible for about 10% of all fish catch in the global ocean. The  $NO_3^-$  and  $O_2$  circulation and budget in this region have a high scientific relevance. This dissertation employs both 2-D box modelling and 3-D coupled physical-biogeochemical modelling to better understand important  $NO_3^-$  and  $O_2$  cycles and budgets in the ETSP.

A simple steady-state box model of a coastal OMZ by Canfield [2006] shows that a positive feedback between  $N_2$  fixation and denitrification could strip the OMZ of all fixed nitrogen when  $N_2$  fixation is permitted to restore the nitrate : phosphate ratio to Redfield proportions in the surface ocean. Observed concentrations of fixed-N (nitrate plus nitrite) in OMZ waters, however, typically range from about 15 to  $40 \mu\text{mol kg}^{-1}$ . We develop a 2-D prognostic box model to examine under which conditions the observed situation of an essentially complete drawdown of subsurface oxygen and an incomplete drawdown of nitrate can be reproduced for the case of the ETSP. The optimum model configuration indicates that the fixed-N inventory can be stabilized at non-zero levels in the ETSP OMZ only if the remineralization rate via denitrification is slower than that via aerobic respiration. For the current notion of nitrogen fixation being favoured in N-deficit waters, the water column of the ETSP could be a small net source of nitrate.

Both atmospheric deposition and benthic remineralization influence the marine nitrogen cycle, and hence ultimately also marine primary production. With the optimum configuration of the 2-D prognostic model, the biological and biogeochemical responses of the ETSP to these processes are analysed. Sensitivity analyses of the local response to both atmospheric deposition and benthic remineralization indicate a nitrogen-balancing mechanism in the ETSP, which tends to keep a balanced nitrogen inventory, e.g., nitrogen input by atmospheric deposition is counteracted by decreasing nitrogen fixation;  $NO_3^-$  loss via benthic denitrification is partly compensated by increased nitrogen fixation; enhanced nitrogen fixation by phosphate regeneration is partly removed by the stronger water-column denitrification. Even though the water column of our model domain acts as a  $NO_3^-$  source, considering benthic denitrification, the ETSP could become a  $NO_3^-$  sink.

Based on the important role of the southern boundary on the oxygen and nutrient contents of

the ETSP found in the box model, a configuration of the same region is built with the high-resolution hydrodynamic model ROMS (*Regional Ocean Modelling System*) [Shchepetkin and McWilliams, 2005], coupled with the biogeochemical model PISCES (*Pelagic Interaction Scheme for Carbon and Ecosystem Studies*) [Aumont and Bopp, 2006]. A validation against in situ observations shows a realistic simulation of the horizontal and vertical oxygen and nutrient distributions by the standard configuration. In the idealised model configurations, designed to investigate the role of southern boundary conditions on the oxygen, nitrate and phosphate variations of the ETSP, the southern boundary is found to have a significant influence on their contents and distributions. Lateral fluxes indicate that our model domain is gaining  $O_2$  and losing  $NO_3^-$  and  $PO_4^{3-}$  through the southern boundary. The southern boundary is found to have a more significant influence on  $O_2$  concentration of the deep layer than the upper layer, suggesting a pathway of deep northward intrusion of well-oxygenated southern waters. The Southern Ocean is thought to be sensitive to global warming due to reductions in deep convection, resulting in significant decrease in  $O_2$  concentrations. This study might give some insights into the impacts of Southern Ocean variations under climate change on the  $O_2$  and nutrient changes of the ETSP.

The mechanisms of the stabilised nitrogen inventory in the OMZ of the ETSP are investigated in the box model, i.e., reduced remineralization rate under suboxic conditions is responsible for the essentially complete drawdown of subsurface oxygen and an incomplete drawdown of nitrate, and the nitrogen fixation and water-column denitrification adjust to counteract the additional nitrogen input via nitrogen deposition and nitrogen removal via benthic denitrification. In the high-resolution coupled physical-biogeochemical model, the stabilised nitrogen inventory is observed in the OMZ of the ETSP even though the remineralisation rates of denitrification and aerobic respiration are identical. Dissolved organic matter (DOM) is found to be exported out of the model domain, which could explain the stabilised nitrogen inventory, because DOM cycling is thought to be capable of promoting the spacial decoupling of nitrogen fixation and denitrification, and allowing for negative feedbacks stabilising the nitrogen inventory [Landolfi et al., 2013].

## Zusammenfassung

Stickstoff ist ein essentieller Nährstoff in der Kontrolle der marinen Primärproduktion. Seine Zirkulation im Ozean ist eng mit primären Zuwachs- und Verlustprozessen wie Stickstofffixierung, atmosphärischer Deposition, Denitrifizierung in der Wassersäule und benthischer Denitrifizierung, und anaerober Ammoniumoxidation (anammox), verbunden. Sauerstoffminimumzonen (OMZs) im Ozean sind für gewöhnlich als Wassersäulen, in denen die  $O_2$ -Konzentration unter  $20 \mu\text{mol kg}^{-1}$  liegt, definiert. Aktuell erstrecken sich OMZs nur über 8 % der globalen Ozeanfläche, sie könnten jedoch für bis zu 30–50 % des gesamten Verlustes an fixiertem Stickstoff an die Atmosphäre, hauptsächlich in Form von  $N_2$ , durch Denitrifizierung oder anammox verantwortlich sein. Aus dem östlichen tropischen Südpazifik (ETSP), wo sich eine der permanenten OMZs im Ozean befindet, stammen über 10% der globalen Fischereierträge.  $NO_3^-$ - und  $O_2$ -Zirkulation und -Haushalt in dieser Region haben hohe wissenschaftliche Relevanz. Diese Dissertation verwendet sowohl 2-D box modelling als auch 3-D gekoppelte physikalisch-biogeochemische Modellierung um wichtige  $NO_3^-$ - und  $O_2$ -Kreisläufe und -Etats im ETSP besser zu verstehen.

Ein einfaches steady-state Box-Modell einer küstennahen OMZ von Canfield [2006] zeigt, dass eine positive Rückkopplung zwischen Stickstofffixierung und Denitrifizierung den gesamten fixierten Stickstoff aus der OMZ herausziehen könnte, wenn es der Stickstofffixierung erlaubt ist, das Nitrat:Phosphat Verhältnis zu Redfield Verhältnissen im Oberflächenozean wiederherzustellen. Beobachtete Konzentrationen fixierten Stickstoffes (Nitrat plus Nitrit) in OMZ-Gewässern erstrecken sich jedoch typischerweise über einen Bereich von 15 bis  $40 \mu\text{mol L}^{-1}$ . Wir entwickeln ein prognostisches 2-D Box-Modell um zu untersuchen, unter welchen Bedingungen die beobachtete Situation einer im Wesentlichen totalen Absenkung des Sauerstoffs unter der Oberfläche und einer unvollständigen Absenkung des Nitrats für den Fall des ETSP reproduziert werden kann. Die optimale Modellkonfiguration weist darauf hin, dass der Etat des fixierten Stickstoffs in der ETSP OMZ nur dann auf von Null verschiedenen Niveaus stabilisiert werden kann, wenn die Remineralisierungsrate durch Denitrifizierung kleiner ist als die durch aerobe Atmung. Nach der aktuellen Auffassung, dass Stickstofffixierung in N-defizitären Gewässern begünstigt wird, könnte die Wassersäule des ETSP eine kleine Nitrat-Nettoquelle sein.

Sowohl atmosphärische Deposition als auch benthische Remineralisierung beeinflussen den marinen Stickstoffkreislauf und damit letztendlich auch die marine Primärproduktion. Mit der optimalen Konfiguration des prognostischen 2-D Modells werden die biologischen und biogeochemischen Reaktionen des ETSP auf diese Prozesse analysiert. Sensitivitätsanalysen der lokalen Reaktion auf atmosphärische Deposition und benthische Remineralisierung zeigen einen stickstoff-ausgleichenden Mechanismus im ETSP auf, der dazu tendiert, den Stickstoffetat im Gleichgewicht zu halten: dem Stickstoffeintrag durch atmosphärische Deposition wird zum Beispiel durch abnehmende Stickstofffixierung entgegengewirkt; der  $NO_3^-$  Verlust durch

benthische Denitrifizierung wird teilweise durch gesteigerte Stickstofffixierung kompensiert; die erhöhte Stickstofffixierung durch Phosphatregeneration wird teilweise durch stärkere Denitrifizierung in der Wassersäule beseitigt. Obwohl die Wassersäule in unserem Modell eine  $\text{NO}_3^-$ -Quelle darstellt, könnte der ETSP, betrachtet man benthische Denitrifizierung, eine  $\text{NO}_3^-$ -Senke sein.

Basierend auf der Entdeckung der wichtigen Rolle des südlichen Ozeans auf den Sauerstoff- und Nährstoffgehalt des ETSP durch das Box-Modell, wird eine Konfiguration der selben Region mit dem hochauflösenden hydrodynamischen Modell ROMS (*Regional Ocean Modelling System*) [Shchepetkin and McWilliams, 2005], gekoppelt mit dem biogeochemischen Modell PISCES (*Pelagic Interaction Scheme for Carbon and Ecosystem Studies*) [Aumont and Bopp, 2006], erstellt. Eine Validierung gegen in situ Beobachtungen zeigt eine realistische Simulation der vertikalen und horizontalen Verteilungen von Sauerstoff und Nährstoffen durch die Standardkonfiguration. In den idealisierten Modellkonfigurationen, die zur Untersuchung der Rolle der südlichen Randbedingungen auf die Sauerstoff-, Nitrat- und Phosphatvariationen des ETSP entwickelt wurden, zeigt sich, dass der südliche Gebietsrand einen signifikanten Einfluss auf deren Gehalt und Verteilungen hat. Die lateralen Flüsse weisen darauf hin, dass unser Modellgebiet  $\text{O}_2$  durch den südlichen Rand erhält und dort  $\text{NO}_3^-$  sowie  $\text{PO}_4^{3-}$  verliert. Es stellt sich heraus, dass der südliche Rand einen signifikanteren Einfluss auf die  $\text{O}_2$ -Konzentrationen in der tieferen Schicht als in der oberen Schicht hat. Dies weist auf einen nordwärts gerichteten Eintragsweg von sauerstoffreichem südlichem Wasser hin. Es wird angenommen, dass der südliche Ozean durch Reduktion in der tiefen Konvektion, die in einer signifikanten Abnahme der  $\text{O}_2$ -Konzentrationen resultiert, empfindlich auf die globale Erwärmung reagiert. Diese Studie könnte Einsichten in die Auswirkungen von Veränderungen im südlichen Ozean durch die zukünftige Klimaveränderung auf die  $\text{O}_2$  und Nährstoffvariationen im ETSP geben.

Die Mechanismen des stabilisierten Stickstoffinventars in der OMZ des ETSP werden im Box-Modell untersucht, dass heißt, die reduzierte Remineralisierungsrate unter suboxischen Bedingungen ist für den im Wesentlichen totalen Verbrauch des Sauerstoffs unter der Oberfläche und den unvollständigen Verbrauch des Nitrats verantwortlich, und die Stickstofffixierung und Denitrifizierung in der Wassersäule regulieren sich dahingehend, den zusätzlichen Stickstoffeintrag via Stickstoffdeposition und Stickstoffentfernung via benthische Denitrifizierung auszugleichen. Im hochauflösenden gekoppelten physikalisch-biogeochemischen Modell wird das stabilisierte Stickstoffinventar in der OMZ des ETSP trotz einer identischen Remineralisierungsrate von Denitrifizierung und aerober Respiration beobachtet. Gelöste organische Stoffe (DOM) werden aus dem Modellgebiet exportiert, wodurch das stabilisierte Stickstoffinventar erklärt werden könnte, da die DOM-Zirkulation als geeignet dafür angesehen wird, die räumliche Entkopplung von Stickstofffixierung und Denitrifizierung zu fördern, und so negative Rückkopplungen, die zur Stabilisierung des Stickstoffinventars führen, zu ermöglichen [Landolfi et al., 2013].

# Chapter 1

## Introduction

### 1.1 The nitrogen cycle in the ocean

#### 1.1.1 The importance of fixed-nitrogen for primary production in the ocean

Nitrogen is an essential component of cells and enzymes responsible for synthetic and metabolic processes of organisms. In the ocean, bioavailable fixed-nitrogen is considered an essential nutrient for marine primary production. Due to the fixed-nitrogen loss via denitrification or anammox in form of  $N_2$  or  $N_2O$  in  $O_2$ -deficit environments, fixed-N is a primary limiting nutrient in the ocean.

#### 1.1.2 Nitrogen species and their transformations in the ocean

In the ocean, nitrogen exists in more chemical forms than most other elements. Most transformations among different species are undertaken by marine organism as part of their metabolism, either to take up nitrogen for organism structure, or to obtain energy for metabolism [Gruber, 2008a]. There are five relatively stable states in the marine environment: the fixed nitrate ( $NO_3^-$ ), nitrite ( $NO_2^-$ ), nitrous oxide ( $N_2O$ ), and ammonium ( $NH_4^+$ ); and unfixed dinitrogen ( $N_2$ ). All of these inorganic species are available for non-diazotrophic phytoplankton except  $N_2$  as the most abundant species, which is only available for diazotrophic phytoplankton. There are also many organic compounds in the ocean containing nitrogen, most of which are in the form of amino-groups.

The main transformations among different species are summarised in Figure. 1.1. In the euphotic zone, where there is enough light to support photosynthesis, the assimilation of  $NH_4^+$  and  $NO_3^-$  to form organic nitrogen performed by marine phytoplankton is the dominant process in the marine nitrogen cycle. This can satisfy their nitrogen demand and fuel the marine

primary production [Gruber, 2008a]. Phytoplankton are thought to preferentially take up  $\text{NH}_4^+$ , because its assimilation is more energy-efficient.  $\text{N}_2$  fixation by diazotrophic phytoplankton is the main process in the ocean to transform  $\text{N}_2$  to bioavailable fixed-Nitrogen.

The organic matter of phytoplankton, which can be transferred to zooplankton and higher-trophic levels in the marine food web, is mainly returned to the environment by remineralization. Under oxic conditions, the organic matter is remineralized to  $\text{NH}_4^+$  initially (named “ammonification”), and then oxidised to  $\text{NO}_3^-$  through nitrification. The remineralization processes for organic nitrogen under anoxic conditions are more diverse and difficult to quantify. Denitrification is thought to be the canonical pathway responsible for anaerobic remineralisation, transferring bioavailable  $\text{NO}_3^-$  to bioavailable  $\text{N}_2$  (Eq. 1.1) [Falkowski, 1997]. Anammox was discovered later as another new pathway responsible for the production of  $\text{N}_2$  in the ocean (Eq. 1.2) [Kuypers et al., 2003].  $\text{N}_2\text{O}$  (not shown in Fig. 1.1), a strong greenhouse gas which is thought to be 200 times more potent than  $\text{CO}_2$ , acts as an intermediary for both nitrification and denitrification, but the relative contributions of these two processes are still debated [Jin and Gruber, 2003].

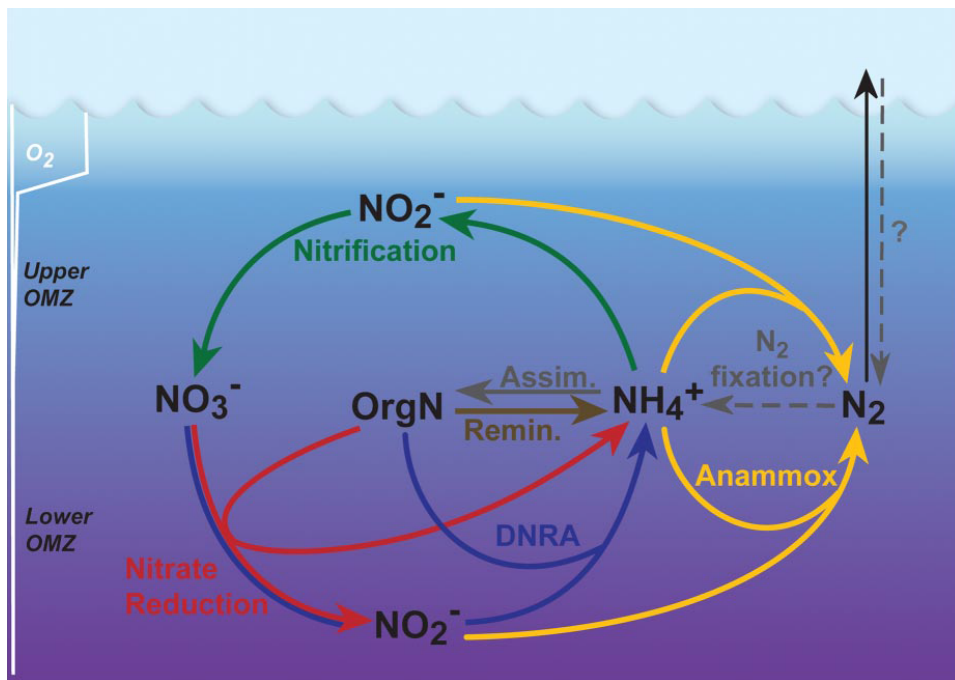
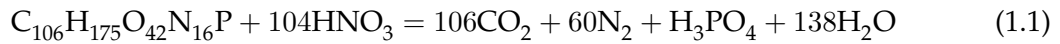


Figure 1.1: Schematic summary of nitrogen cycle in the ocean [Lam et al., 2009].

### 1.1.3 Distribution of fixed-nitrogen in the ocean

#### Global scale

The distribution of different fixed-nitrogen species in the ocean is mainly connected to the biological processes and the physical circulation. Many data exist for profiles of  $\text{NO}_3^-$  and  $\text{NO}_2^-$ , but only sparse data for  $\text{NH}_4^+$ .

The vertical profile of  $\text{NO}_3^-$  follows a prevalent pattern of depletion at the surface and repletion in the deep ocean, which is due to biological uptake by phytoplankton in the surface ocean and remineralization at depth (left graph of Fig. 1.2). Different from  $\text{NO}_3^-$ ,  $\text{NO}_2^-$  and  $\text{NH}_4^+$  concentrations reach maxima at about 50 to 80 meter (lower or bottom part of the euphotic zone), and then decrease rapidly below (left graph of Fig. 1.2). For  $\text{NO}_2^-$ , the maximum occurs at the bottom of euphotic zone or below (right graph of Fig. 1.2), where light is low enough to prevent nitrification limitation and  $\text{NO}_2^-$  is produced during nitrification. The  $\text{NH}_4^+$  maximum is at the deeper part of euphotic zone where there is rapid recycling of organic matter (OM) via remineralization, liberating much of  $\text{NH}_4^+$  (right graph of Fig. 1.2). At the lower part of euphotic zone, photosynthesis may be limited partly by irradiance making generation of  $\text{NH}_4^+$  exceed its consumption.

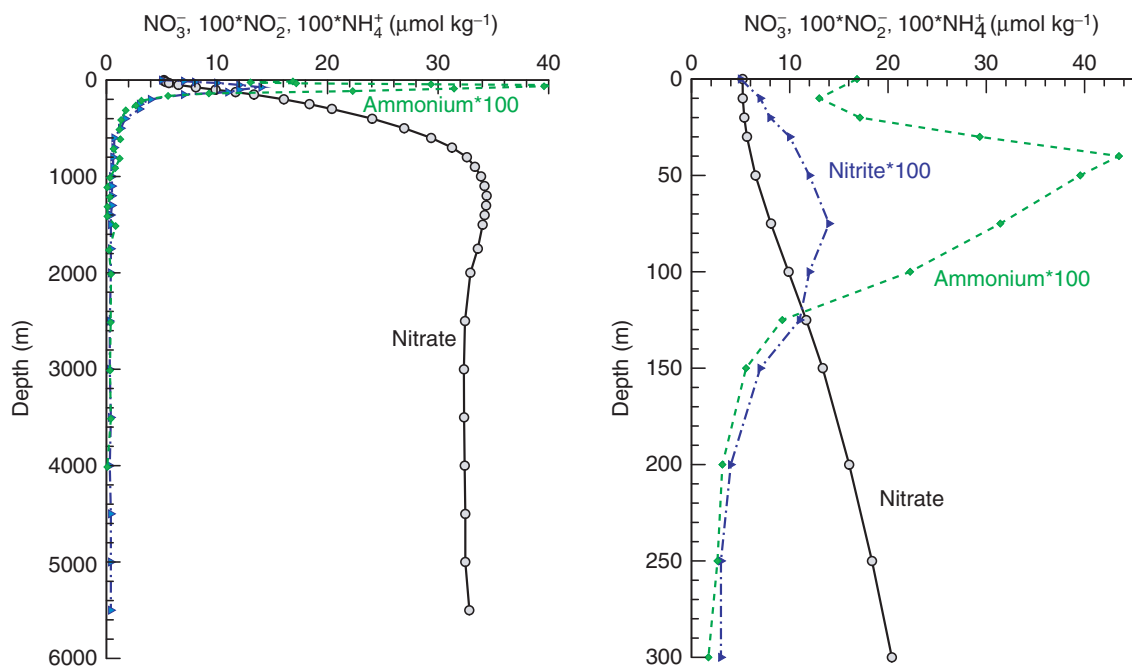


Figure 1.2: Global mean profile of fixed-nitrogen for the entire water volume and top 300 meter [Gruber, 2008a].

The  $\text{NO}_3^-$  concentration in the surface ocean is extremely low except in the low-latitude upwelling regions, including the Southern Ocean, part of the North Pacific and the equatorial

Pacific. Nitrate, together with other nutrients, is upwelled into the surface ocean in the upwelling regions, which can support high primary production. However, even though there is enough fixed-nitrogen in the surface ocean of the three regions mentioned above, there are still low-chlorophyll concentrations there. These are referred as High-Nutrients Low-Chlorophyll (HNLC) regions. The micronutrient iron is thought to be the main limiting factor in HNLC regions [Aumont and Bopp, 2006, Okin et al., 2011].  $\text{NO}_2^-$  and  $\text{NH}_4^+$  concentrations in the ocean are almost three orders of magnitude lower than that of  $\text{NO}_3^-$ . Thus, we mainly focus on  $\text{NO}_3^-$  in our work.

### Eastern Tropical South Pacific (ETSP)

The ETSP hosts one of the major permanent OMZs in the ocean. Due to the high rate of organic matter decomposition and insufficient ventilation, the oxygen minimum zone ensues, concomitant with fixed-N loss in form of  $\text{N}_2$  or  $\text{N}_2\text{O}$  by denitrification or anammox. Even though there is fixed-N loss in the  $\text{O}_2$ -deficit region, the  $\text{NO}_3^-$  concentration in the ETSP ranges roughly from 15 to 45  $\mu\text{mol kg}^{-1}$  [Codispoti and Christensen, 1985, Chang et al., 2010].

Usually,  $\text{NH}_4^+$  concentrations are almost undetectable in coastal upwelling regions except in some parts of the lower euphotic zone and the bottom water of the continental shelf. Uniquely, there is a relatively large amount of  $\text{NH}_4^+$  provided by excretion from anchovy in the upwelling region off Peru [Gruber, 2008b]. The relative high concentration of  $\text{NH}_4^+$  can also be caused by the dissimilatory nitrate reduction to ammonium (DNRA) in the sediment [Bohlen et al., 2012].

#### 1.1.4 Fixed-N budget in the ocean

Nitrogen is thought to be the proximate limiting nutrient for growth of phytoplankton, thus, the nitrogen budget and its spatio-temporal distribution are crucial for marine primary production. The fixed-N inventory in the ocean is determined by the differences between sources and sinks. Due to the large-scale of the different processes and lack of enough measurements, various biogeochemical models coupled with hydrodynamic models are employed to estimate the quantity of different sources and sinks of fixed-nitrogen besides in-situ observations. Here we outline the most known sinks and sources estimated from both models and observations.

### Sources

#### *N<sub>2</sub> fixation*

In the last decades, tremendous progress has been made in quantifying  $\text{N}_2$  fixation rates and organisms responsible for fixing atmospheric  $\text{N}_2$  both in sediments and the pelagic ocean [Carpenter, 1983, Capone et al., 1997, Montoya et al., 2004]. Macrophyte communities, microbial



mats and coral reef associated habits are found to be responsible for benthic nitrogen fixation. In the water column, early research mainly focused on cyanobacterium *Trichodesmium* [Karl et al., 2002], however, subsequent discoveries have shown that unicellular coccoid cyanobacteria, and certain diatom- or dinoflagellate-symbiont cyanobacteria can also express nitrogenase, enriching the diversity of  $N_2$  fixers [Carpenter and Capone, 2008, Montoya et al., 2004, Foster and Zehr, 2006]. Since oceanic nitrogen fixation is also affected by temperature, light, oxygen, turbulence, trace metals and inorganic nutrients (e.g. phosphate), estimating its rate is a huge challenge.

Due to the large spatial and temporal variability and continuing discoveries of new species responsible for  $N_2$  fixation, there are still considerable uncertainties in the estimates for the global and large-scale regional rates of  $N_2$  fixation. The estimated global rates have increased from about  $5 \text{ Tg N yr}^{-1}$  to about  $300 \text{ Tg N yr}^{-1}$  [Capone and Budin, 1982, Brandes and Devol, 2002]. Most recent biogeochemical model results indicate a range of  $100\text{--}300 \text{ Tg N yr}^{-1}$ , which results from insufficiently constrained biogeochemical parameters [Codispoti et al., 2001, Gruber, 2004, Deutsch et al., 2007, DeVries et al., 2012]. It is also difficult to estimate the  $N_2$  fixation rate in upwelling regions such as the ETSP, the ETNP and the Arabian Sea. New measurement methods and discoveries of new species for  $N_2$  fixation will probably allow better constraints for the oceanic  $N_2$ -fixation rate [Foster et al., 2011, Großkopf et al., 2012].

#### *Atmospheric nitrogen deposition*

In the last few decades, a number of investigations have been carried out on the role of atmospheric fixed-nitrogen input into the ocean on marine biogeochemical cycles [Duce, 1986, Duce et al., 1991, Krishnamurthy et al., 2007, 2010, Okin et al., 2011]. Duce et al. [2008] indicate that the anthropogenic nitrogen deposition input is rapidly approaching the global oceanic estimates for  $N_2$  fixation, arising from an order of magnitude lower than  $N_2$  fixation at the pre-industrial stage. However, the response of nitrogen-fixation and denitrification to atmospheric nitrogen deposition remains an open question. Inorganic nitrogen, from fossil fuel combustion, biomass burning, soil emission and breakdown of urea from domestic animals [Warneck, 1988], dominates atmospheric nitrogen inputs into the global ocean [Paerl and Whitall, 1999]. The species of nitrogen from deposition of Lamarque et al. [2011] only include inorganic  $\text{NO}_y$  and  $\text{NH}_x$ . The magnitude of dissolved organic nitrogen (DON) deposition is not clear due to lack of observations [Cornella et al., 2003, Duce et al., 2008, Zamora et al., 2011]. Even though phytoplankton can take up DON [Seitzinger and Sanders, 1999, Duarte et al., 2006, Duce et al., 2008], the contribution of DON to total nitrogen deposition is still debated [Duce et al., 2008, Zamora et al., 2009] and the distribution, bioavailability and lifetime are also not clear. Therefore, including DON deposition in the analysis is still challenging.

#### *Riverine discharge*

A significant quantity of nitrogen from land, mainly from fertilizer use, fossil fuel combustion and biological nitrogen fixation, is transported from the continents via rivers into the coastal

oceans [Boyer et al., 2006]. The riverine nitrogen discharge into the ocean includes dissolved inorganic nitrogen (DIN), dissolved organic nitrogen (DON) and particulate nitrogen (PN). There is a large uncertainty in quantifying riverine nitrogen input into the ocean at global and regional scales due to the coarse resolution of available data, land characteristics and associated assumptions about their hydraulic characteristics, vague nitrogen transportation processes and insufficient measurements for riverine nitrogen loads [Boyer et al., 2006]. Based on current comprehensive assumptions and empirical models, the global riverine nitrogen input is estimated from 40 to 66 Tg N yr<sup>-1</sup> [Green et al., 2004, Seitzinger et al., 2005].

The integrated model of Seitzinger et al. [2010] indicates that south Asia is a region responsible for major riverine nitrogen export to the coastal ocean, however, south America together with Africa show the largest increases in DIN input in the past 30 years and in the coming 30 years. The ETSP will probably receive more riverine nitrogen input in the future and riverine nitrogen input will account for more contribution to nitrogen sources in ETSP.

## Sinks

In the global marine environment, water-column denitrification and anammox in OMZs, together with benthic denitrification, determine the magnitude of fixed-nitrogen loss, in which benthic denitrification is considered a major fixed-nitrogen loss process [Codispoti et al., 2001, Galloway et al., 2004, Gruber, 2004, Codispoti, 2007, Somes et al., 2013].

### *Water-column denitrification and anammox*

OMZs play an important role in the global marine fixed-nitrogen budget as they are sites responsible for a large fraction of total marine fixed-N loss [Canfield, 2006]. The relative contribution of heterotrophic denitrification and autotrophic anammox to the total oceanic fixed-nitrogen sink remains debated [Kuypers et al., 2005, Lam et al., 2009, Ward et al., 2009]. Even though autotrophic anammox has been suggested to be responsible for most of the fixed-N loss in the ETSP [Lam et al., 2009, Kalvelage et al., 2013], anammox is ultimately driven by the flux of organic matter into the OMZ as heterotrophic remineralization of organic matter supplies the necessary substrates for it [Koeve and Kähler, 2010, Kalvelage et al., 2013]. Thus, heterotrophic denitrification is considered as the major fixed-nitrogen loss process in the present study.

Whether canonical denitrification or anammox is dominant in marine fixed-nitrogen loss still remains uncertain, since they require and happen in the same environmental conditions. Most estimates for total N-loss in the water column based on models and in-situ observations lie between 50–150 Tg N yr<sup>-1</sup> [Gruber, 2004, Codispoti, 2007].

### *Benthic denitrification*

A high rain rate to the sediment can drive microorganisms to use up all  $O_2$  and result in an anaerobic sediment environment, which provides conditions for fixed-nitrogen loss. Continental shelves and the upper continental slopes are thought to be the most important sites for benthic fixed-nitrogen loss [Christensen et al., 1987, Devol, 1991]. Bohlen et al. [2011] found that the anaerobic continental shelf and upper continental slope of the ETSP across a section at  $11^\circ S$  are sites of fixed-nitrogen recycling rather than nitrogen loss, because of relatively low rates of denitrification and high rates of  $NH_4^+$  release from Dissimilatory Nitrate Reduction to Ammonium (DNRA). Besides, the contribution of open-ocean sediments is still uncertain. Because of the large uncertainties, the global benthic denitrification rate is estimated in a large range from about 60 [Codispoti and Christensen, 1985] to more than 300  $Tg\ N\ yr^{-1}$  [Codispoti, 2007].

### 1.1.5 The sensitivity of the marine nitrogen-cycle to phosphate in the ocean

Compared with the nitrogen cycle, the phosphorus cycle in the ocean is relatively simple, because P has no gas phase and only one valence (+5) in its compound forms. The main global sources of P into the ocean are thought to be river runoff, atmospheric deposition and volcanic sources, among which river runoff is thought to be most significant. Sedimentary burial is the dominant loss pathway for P. In the sediment under anoxic conditions, P regeneration provides extra inorganic P release into the water column from the sediment [Wallmann, 2010].

Both N and P are essential macronutrients for growth of phytoplankton in the ocean. According to the classical Redfield ratio, the N:P stoichiometry 16 is regarded as the criteria to differentiate between N limitation and P limitation in the ocean [Redfield, 1934]. A ratio less than 16 is assumed to indicate N limitation, and a ratio larger than 16 is assumed to indicate P limitation. There are debates about which nutrient, P or N, controls primary production in the ocean. The geological viewpoints support that P is the limiting nutrient because N can be supplied by nitrogen fixation when it limits the growth of phytoplankton and the residence time of P is much longer than that of N in the ocean [Ruttenberg, 2003]. Fixed-nitrogen is thought to be the limiting nutrient by biologists because there is a relatively low N:P ratio in many surface ocean regions and nitrogen enrichment experiments do enhance primary production there [Falkowski et al., 1998, Garcia et al., 2010b]. Tyrrell [1999] uses a model to indicate that N is the proximate limiting nutrient in the ocean, while P is considered as the ultimate limiting nutrients for primary production. There is evidence that P is the limiting nutrient for nitrogen fixation in the North Atlantic and North Pacific [Karl et al., 1997, Wu et al., 2000], thus, the addition of P into these regions could enhance the biological fixed-nitrogen input into ocean.

P regeneration has been found on the continental shelf off California and off the Peruvian coast, where anoxic bottom water enhances P release from sediments receiving a high rain rate [Ingall and Jahnke, 1994]. The enhanced release of P from the sediment can also enhance primary

production, which could in turn stimulate oxygen depletion in bottom waters significantly, thereby providing further P addition for primary production [Ingall et al., 1993]. The intensification of anoxic conditions in the coastal ocean due to anoxic P regeneration could also have significant influence on the nitrogen cycle in these regions.

## 1.2 Oxygen minimum zones (OMZs) in the ocean

### 1.2.1 Definition and Characteristics

Since the expression of “OMZ” appeared in Cline and Richards [1972], Oxygen minimum zones (OMZs), characterized as  $O_2$ -deficit water columns, receive much attention in the present ocean research. Commonly, an OMZ is defined as water with  $O_2$  concentration less than  $20 \mu\text{mol kg}^{-1}$ , but this threshold is still a matter of debate [Paulmier and Ruiz-Pino, 2009, references therein]. For example, it has been suggested that denitrification, which is  $O_2$ -dependent, can be the criterion for defining the extent of an OMZ [Paulmier and Ruiz-Pino, 2009].

### 1.2.2 OMZ distribution in the ocean

Using the global World Ocean Atlas 2005 (WOA2005)  $O_2$  climatology, four major tropical OMZs in the open ocean have been described by Paulmier and Ruiz-Pino [2009]: the eastern tropical South Pacific (ETSP) and eastern tropical North Pacific (ETNP), in the Pacific Ocean; the Arabian Sea (AS) and Bay of Bengal (BB), in the Indian Ocean (Fig. 1.3). More recently, the eastern sub-tropical North Pacific ( $25 - 52^\circ N$ ) has been identified as another permanent deep OMZ. Two additional seasonal OMZs at high latitudes have also been identified: the western Bering Sea and the Gulf of Alaska. The total ocean area and volume occupied by the most intense OMZs ( $O_2 < 20 \mu\text{mol kg}^{-1}$ ) have been evaluated by WOA 2005 data analysis [Paulmier and Ruiz-Pino, 2009]:  $30.4 \pm 3$  million  $\text{km}^2$  and  $102 \pm 15$  million  $\text{km}^3$ , accounting for, 8% and 1% of the global ocean area and volume respectively.

Paulmier et al. [2006] demonstrate that there are at least three layers in the structure of OMZs when analyzing the OMZ in the ETSP off Chile: the oxycline with an extremely high  $O_2$  gradient, the core with  $O_2$  concentration less than  $20 \mu\text{mol kg}^{-1}$ , and a deep  $O_2$  gradient. The oxycline is thought to be the region where most intensive remineralization occurs [Brandes et al., 2007]. The structure of the OMZ-core is diverse. The core of the OMZ in the eastern tropical Pacific is thought to lie between 150 and 400 meter, with an oxygen minimum of  $0.1 \mu\text{mol kg}^{-1}$  [Karstensen et al., 2008]. The OMZ core of the Indian Ocean corresponds to depths between 100 and 650 meter and reaches  $O_2$  values as low as  $0.3 \mu\text{mol kg}^{-1}$  [Morrison et al.,

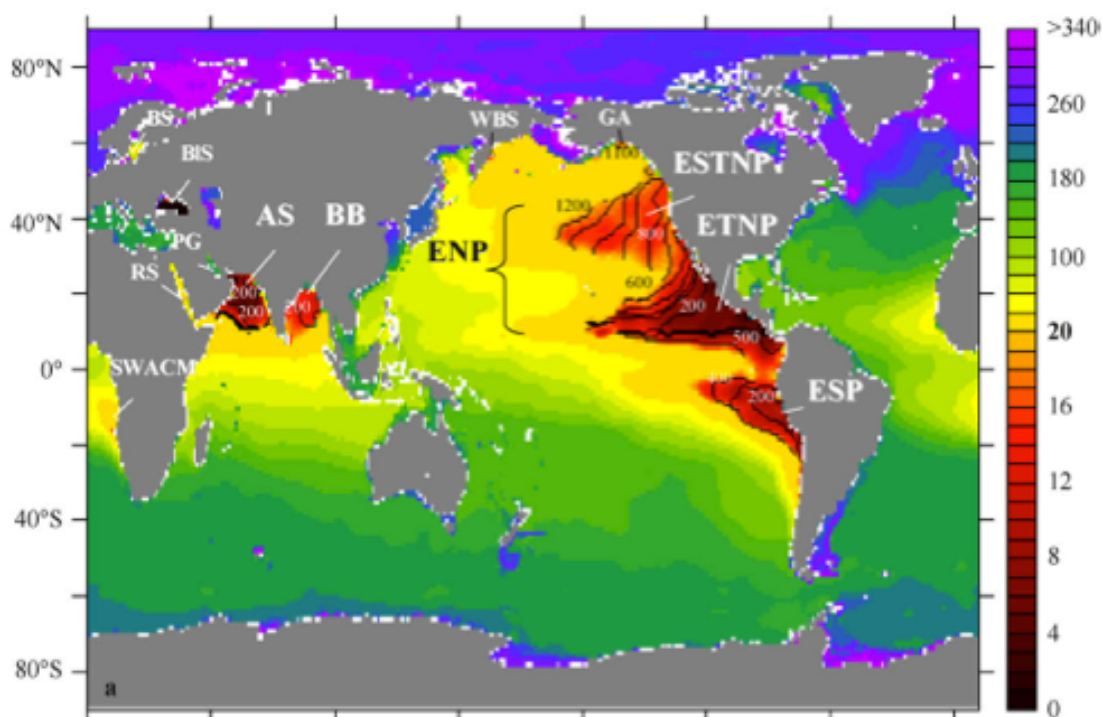


Figure 1.3: OMZs distribution in the ocean [Paulmier and Ruiz-Pino, 2009].

1999]. There is no suboxic zone ( $<20 \mu\text{mol kg}^{-1}$ ) in Atlantic Ocean, where the  $\text{O}_2$  concentration minimum corresponds to a depth of about 420 and 400 meter, respectively, in the North and South Atlantic ( $\text{O}_2 > 40 \mu\text{mol kg}^{-1}$  in the north;  $>20 \mu\text{mol kg}^{-1}$  in the south, except in some areas over the continental margin where  $\text{O}_2$  reaches 17–18  $\mu\text{mol kg}^{-1}$ ).

### 1.2.3 Importance of OMZs and their significance for the marine nitrogen cycle

The intensity of perturbations of OMZs and their potential influence on the marine biogeochemical cycle and ecosystems depend on their extent, which can vary in response to climate change, such as lower ventilation due to stratification and decrease of  $\text{O}_2$  solubility at higher temperature, and natural or anthropogenic perturbations through limiting nutrient inputs by upwelling, river input or atmospheric dust deposition [Stramma et al., 2008]. These are the keys to understanding the present unbalanced or balanced nitrogen cycle and the role of the ocean in sequestering atmospheric greenhouse gases [Codispoti, 2007, DeVries et al., 2012], since the  $\text{O}_2$ -deficit in the marine environment is essential for the anaerobic bacteria which can convert fixed-N to  $\text{N}_2$  or  $\text{N}_2\text{O}$  via denitrification or anaerobic ammonium oxidation (anammox) [Devol, 2003, Kuypers et al., 2003]. Under  $\text{O}_2$ -deficit conditions,  $\text{NO}_3^-$  can replace  $\text{O}_2$  as electron acceptor in decomposition of organic matter in denitrification, in which  $\text{NO}_3^-$  is reduced to  $\text{N}_2$  [Devol, 2003]. The bacteria responsible for anammox (e.g., bacteria from the Planctomycetales

order) can oxidize  $\text{NH}_4^+$  directly to  $\text{N}_2$  with  $\text{NO}_2^-$  as the electron acceptor [Kuypers et al., 2003].

#### 1.2.4 The oxygen minimum zone of the ETSP

The Humboldt Current System is the most productive eastern boundary current system. This work focuses on the northern part of the Humboldt Current System, the ETSP off Peru, hosting one of the most significant permanent OMZs. The ETSP upwelling region is known as one of the most productive systems of the world ocean, where high export production is decomposed in the water column below, consuming dissolved oxygen. Combined with sluggish ventilation, the high  $\text{O}_2$  demand produces the sub-surface OMZ of the ETSP [Hellya and Levin, 2004]. In oxygen-depleted regions,  $\text{NO}_3^-$  is converted to  $\text{N}_2\text{O}$  or  $\text{N}_2$  by canonical denitrification or anammox, thus resulting in a nitrate deficit relative to the Redfield equivalent of phosphate [Codispoti and Christensen, 1985].

Figure 1.4 is a schematic summary of the current systems in the eastern tropical Pacific. Due to the remarkable characteristic of the Peru Current System (PCS) being in proximity and directly connected to the equatorial ocean, the Equatorial Current System (ECS) can not be excluded when the PCS is discussed.

The surface of the PCS is dominated by the Peru Coastal Current (PCC), which is equatorward, with a width of 500-1000 km from the coast [Strub et al., 1998]. The PCC is directly associated with coastal upwelling, thus featuring cold and salty water. It feeds the South Equatorial Current (SEC), which is the westward surface current of the ECS. There is a poleward flow below the PCC dominating the subsurface layer (Fig. 1.4). The Peru-Chile Under Current (PCUC) can extend from 50m to 700m along the western coast of South American with depth increasing toward the south, transporting salty water rich in nutrients and extremely low in  $\text{O}_2$  [Wyrski, 1963, Penven et al., 2005, and references therein]. The PCUC is thought to originate from the eastward Equatorial Under Current (EUC), which splits when arriving at the American landmass. Both the primary South Subsurface Countercurrent (pSSCC) and secondary South Subsurface Countercurrent (sSSCC) could be complementary sources of the PCUC [Lukas, 1986, Montes et al., 2010]. Since the PCUC is thought to be one of the major sources of the upwelled water in the ETSP, its oxygen and nutrient contents are important for the study of the upwelling region and the OMZ below [Penven et al., 2005, and references therein]. The connection between the PCUC and the equatorial eastward subsurface currents (EUC, pSSCC, sSSCC), especially with respect to oxygen and nutrients, is very important for understanding the formation of the OMZ in the ETSP. However, Montes et al. [2010] indicate in a high-resolution regional model that only about 30% of the PCUC is fed by the three subsurface equatorial currents (EUC, pSSCC, sSSCC), among which only the two SSCC contribute substantially. The remaining part of the PCUC comes from an alongshore recirculation associated with flows below and from the southern part of the domain, south of  $9^\circ\text{S}$  [Montes et al.,

2010].

Another surface current belonging to the PCS, the offshore Peru Oceanic Current (POC) extends from the surface to a depth of 700m and veers eastward around 15–20°S [Wyrski, 1963, Penven et al., 2005]. Offshore of the PCC and the PCUC, the Peru-Chile Counter Current (PCCC) is featured by warm water originating from the split of EUC, and ranges between 80°W–84°W and 8°S–35°S [Huyer et al., 1991, Penven et al., 2005]. The PCCC flows southward and reaches a maximum speed of  $0.1\text{m s}^{-1}$  at 50m depth [Huyer et al., 1991]. So far, the connection and exchanges of nutrient and oxygen between the PCCC and the PCUC are not clear.

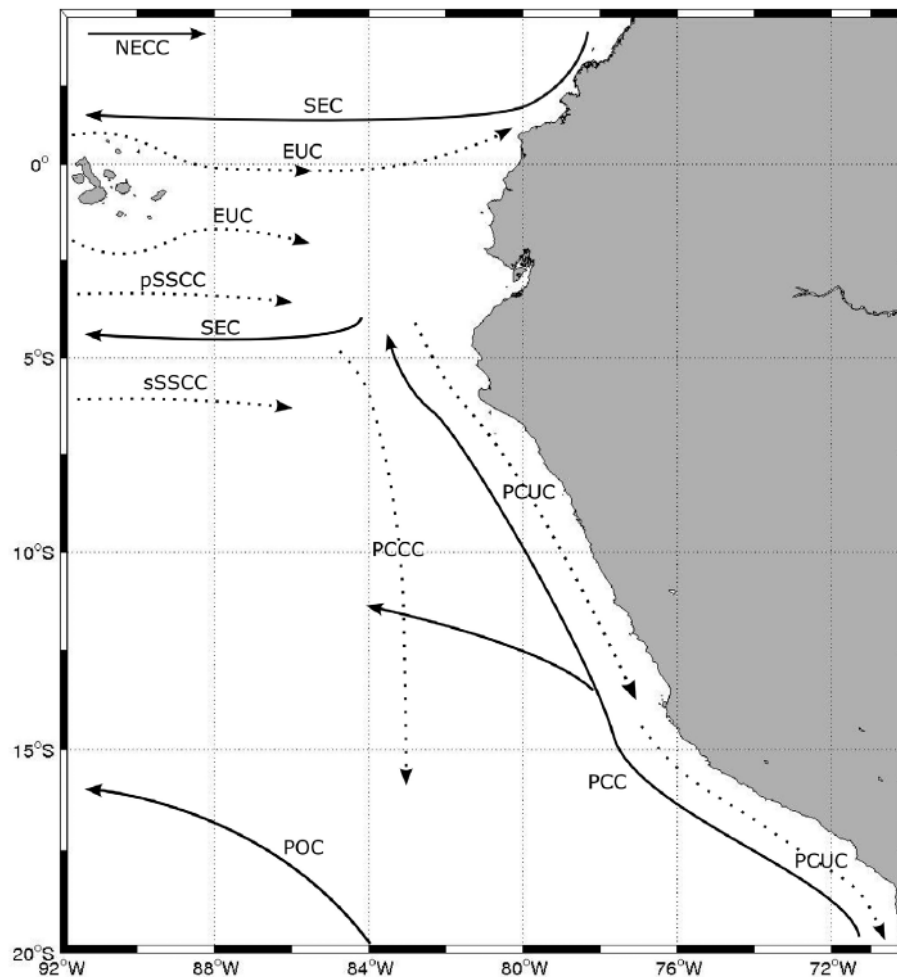


Figure 1.4: Oceanic circulation scheme for the eastern tropical Pacific [Montes et al., 2010, and references therein]. Solid lines indicate the surface currents and dotted lines represent subsurface currents. NECC: North Equatorial Counter Current; SEC: South Equatorial Current; EUC: Equatorial Undercurrent; pSSCC: primary South Subsurface Counter Current; sSSCC: secondary Southern Subsurface Counter Current; PCC: Peru Coastal Current; POC: Peru Oceanic Current; PCUC: Peru-Chile Undercurrent; and PCCC: Peru-Chile Counter Current.

### 1.3 Models

Two modelling approaches are applied in this dissertation to explore the nitrogen cycle in the OMZ of the ETSP: a fully prognostic 5-box model coupled with a simplified biogeochemical model and the 3-D dynamical regional model ROMS (*Regional Ocean Modelling System*) coupled with PISCES (*Pelagic Interaction Scheme for Carbon and Ecosystem Studies*).

#### 1.3.1 Box model

Even though there are lots of state-of-the-art global circulation models (GCM) available, box models are still popular tools applied to study global and regional marine biogeochemical cycles [Shaffer and Sarmiento, 1995, Tyrrell, 1999, Deutsch et al., 2004, Canfield, 2006, Mills and Arrigo, 2010, Eugster and Gruber, 2012, DeVries et al., 2012], due to their ability to capture the most important first-order characteristics of the system and computational efficiency required for thousands of sensitivity or calibration experiments. Recently, a dynamic 5-box model of the ETSP was presented by Mills and Arrigo [2010] to explore the impact of stoichiometry of dissolved nitrogen and phosphorus utilization by phytoplankton on the magnitude and distribution of nitrogen fixation. Eugster and Gruber [2012] used the 14-box model of Deutsch et al. [2004] to estimate the global marine nitrogen budget mainly determined by the magnitude of  $N_2$ -fixation and denitrification.

We developed Canfield's [2006] box model into a fully prognostic box model of  $NO_3^-$ ,  $PO_4^{3-}$  and  $O_2$  cycles in a coastal upwelling region and an adjacent ocean basin. Different from Canfield's [2006] model, which includes only the dynamics of the OMZ with all the biogeochemical tracers prescribed in the surrounding waters, our box model represents local as well as spatially-separated feedbacks between the relatively small OMZ and the much larger open-ocean basin. The physical dynamics of our model system is governed by vertical and horizontal mixing and advection, which are calibrated with  $\Delta^{14}C$  data from GLODAP [Key et al., 2004].

A simplified biogeochemical model is incorporated, including two phytoplankton species: ordinary phytoplankton, which require both  $NO_3^-$  and  $PO_4^{3-}$  for growth, and nitrogen fixers, which can grow only if  $PO_4^{3-}$  is available in the ambient environment. There are complex N-cycle biogeochemical models with increasing numbers of compartments developed [Laws et al., 2000, Fennel et al., 2006]. But there are still some suggestions that modellers should stick with simple NPZ and NPZD model, rather than try to 'parameterize' additional food web complexity [Steele, 1998, Pahlow and Vézina, 2003]. In the box model, we keep our biogeochemical model as simple as possible to match the simple physical dynamics.



### 1.3.2 3-D model

The 3-D model employed in this dissertation is the Regional Ocean Modelling System (ROMS-AGRIF) [Shchepetkin and McWilliams, 2005] coupled with the biogeochemical model PISCES [Aumont and Bopp, 2006].

ROMS is a new generation ocean circulation model that has been specially designed for accurate simulations of regional systems. A split-explicit, free surface, terrain-following vertical coordinates ocean model which solves the primitive equations of Navier Stokes in the presence of rotation characterises ROMS [Shchepetkin and McWilliams, 2005]. Different from many other oceanic general circulation models, ROMS has sigma vertical coordinate, which is designed to enhance resolution near the sea surface and employs a mode-splitting algorithm to separate the natural time-scale between barotropic and baroclinic processes, by solving the vertically-integrated barotropic momentum equations [Shchepetkin and McWilliams, 2009, Penven et al., 2010].

This work requires a biogeochemical model which can simulate the  $O_2$ ,  $NO_3^-$  and  $PO_4^{3-}$  cycles of the ETSP, include the explicit processes for the marine nitrogen cycle in the OMZ (denitrification), and also nitrogen-fixation. PISCES, derived from the Hamburg Model of Carbon Cycle (version 5, HAMOCC5), is adequate to reproduce the biogeochemical processes required for the simulation of the ETSP.

PISCES is a biogeochemical model which simulates marine biogeochemical productivity and describes the biogeochemical cycles of carbon and the main nutrients (P, N, Si, Fe) [Aumont and Bopp, 2006]. It includes two phytoplankton types, which correspond to nanophytoplankton and diatoms, and two zooplankton size classes that represent microzooplankton and mesozooplankton. The limiting nutrients for phytoplankton growth include:  $NO_3^-$  and  $NH_4^+$ ,  $PO_4^{3-}$ , Si and Fe. There are three non-living compartments: semi-labile dissolved organic matter, small and big sinking particles. The model also simulates dissolved inorganic carbon, total alkalinity and dissolved oxygen [Aumont et al., 2003]. All the parameters for this region are according to Echevin et al. [2014].

The biogeochemical processes related to the nitrogen cycle include nitrogen fixation, denitrification, nitrification and ammonification in PISCES [Aumont and Bopp, 2006]. Diazotrophic phytoplankton can fix  $N_2$  under the following conditions: 1) water temperature is above  $20^\circ C$ , 2) there is insufficient fixed-N in the environment, 3) iron is available. Nitrogen fixation is only restricted to the sea surface and annual total nitrogen fixation should balance denitrification to ensure nitrogen conservation in the model domain. Denitrification ensues when oxygen concentration is under  $6 \mu mol kg^{-1}$ . Nitrification represents the conversion of  $NH_4^+$  to  $NO_3^-$  due to bacterial activity, which is assumed to be photoinhibited and  $O_2$  concentration dependent. Ammonification is responsible for the remineralization of particulate organic matter and dissolved organic matter to  $NH_4^+$ .

## 1.4 Overview of the thesis and author contributions

Nitrogen is an essential nutrient for phytoplankton in the ocean, thus, the bioavailable fixed-nitrogen budget is an essential control on the potential of the ocean to sequester atmospheric CO<sub>2</sub> via the marine biological pump. The nitrogen budget in the ocean is determined by the main sources, i.e., nitrogen fixation and atmospheric deposition, and sinks, including denitrification and anammox. O<sub>2</sub> concentration is an important control on the marine nitrogen cycle, because NO<sub>3</sub><sup>-</sup> is the second electron acceptor after O<sub>2</sub> for the remineralization of organic matter. In suboxic conditions, biological fixed-nitrogen is lost via denitrification or anammox. Due to the current climate warming, O<sub>2</sub> concentration in the ocean is thought to be decreasing due to lower solubility and intensified stratification, which could also have an impact on the marine nitrogen cycle. This dissertation concentrates mainly on the nitrogen and O<sub>2</sub> cycles of the eastern tropical South Pacific and its OMZ.

Chapter 2 examines under which conditions the observed situation of an essentially complete drawdown of subsurface oxygen and an incomplete drawdown of nitrate can be reproduced for the case of the ETSP OMZ. It reveals that the fixed-N inventory can be stabilized at non-zero levels only if the remineralization rate via denitrification is slower than that via aerobic respiration. This conclusion is drawn by developing a fully prognostic box model of NO<sub>3</sub><sup>-</sup>, PO<sub>4</sub><sup>3-</sup> and O<sub>2</sub> cycles in the coastal upwelling of the ETSP and its adjacent ocean basin. Model results also demonstrate that the water column of the ETSP is a NO<sub>3</sub><sup>-</sup> source. This chapter is published in the journal *Biogeosciences* (citation: B. Su, M. Pahlow, H. Wagner, and A. Oschlies, What prevents nitrogen depletion in the oxygen minimum zone of the eastern tropical South Pacific?, *Biogeosciences*, 12, 1113–1130, doi:10.5194/bg-12-1113-2015). BS developed the fully dynamic 5-box prognostic model from the Canfield [2006] model based on the initial idea provided by AO. BS performed all experiments, calculations and analysis with the help of MP. BS wrote the manuscript, with guidance and comments from all co-authors.

Chapter 3 investigates the sensitivity of the nitrogen cycle in the ETSP to nitrogen deposition and benthic remineralization, employing the optimum configuration developed by Su et al. [2015]. The local response to both atmospheric deposition and benthic remineralization is examined in the sensitivity analyses. The results indicate a nitrogen-balancing mechanism in the ETSP, which tends to keep a balanced nitrogen inventory, i.e., nitrogen input by atmospheric deposition is counteracted by decreasing nitrogen fixation; NO<sub>3</sub><sup>-</sup> loss via benthic denitrification is partly compensated by increased nitrogen fixation; enhanced nitrogen fixation by phosphate regeneration is partly removed by the stronger water-column denitrification. Even though the water column in our model domain acts as a NO<sub>3</sub><sup>-</sup> source, the ETSP including benthic denitrification might become a NO<sub>3</sub><sup>-</sup> sink. This chapter is a submitted manuscript entitled “Box-modelling the impacts of atmospheric nitrogen deposition and benthic remineralization on the nitrogen cycle of the eastern tropical South Pacific” by B. Su, M. Pahlow and A. Oschlies. BS performed all the data analysis, calculation and sensitivity experiments based on the initial

idea provided by AO. BS wrote the manuscript with the guidance and comments from MP and AO.

In chapter 4, a configuration of the ETSP OMZ is built employing the 3-D hydrodynamic model ROMS coupled with the biogeochemical model PISCES. Three experimental model configurations are performed to investigate the influence of southern boundary conditions on the  $O_2$  and nutrient distributions of the model domain. This chapter is a manuscript in preparation with the title "Coupled physical/biogeochemical modeling of the oxygen minimum zone of the eastern tropical South Pacific: sensitivity to the southern boundary conditions" by B. Su, Y. José, I. Montes, M. Pahlow, and A. Oschlies. BS conceived the idea and designed the experiments with the guidance of AO and IM. BS performed all experiments, calculations and analyses with help of YJ and MP. BS wrote the manuscript with comments and english improvement provided by the co-authors.

In Chapter 5, the main results of the thesis are summarised. The mechanism responsible for the stabilised nitrogen inventory in the ROMS-PISCES model and the influence of interannual variabilities (i.e., ENSO) on the dynamics of the OMZ and the nitrogen balance will be investigated as outlook on future research. To have a better understanding of the full nitrogen cycle in the ETSP, coupling the benthic remineralization into ROMS-PISCES will be another future research direction.



## Chapter 2

# What prevents nitrogen depletion in the oxygen minimum zone of the eastern tropical South Pacific?

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**Abstract** Local coupling between nitrogen fixation and denitrification in current biogeochemical models could result in runaway feedback in open-ocean oxygen minimum zones (OMZs), eventually stripping OMZ waters of all fixed nitrogen. This feedback does not seem to operate at full strength in the ocean, as nitrate does not generally become depleted in open-ocean OMZs. To explore in detail the possible mechanisms that prevent nitrogen depletion in the OMZ of the eastern tropical South Pacific (ETSP), we develop a box model with fully prognostic cycles of carbon, nutrients and oxygen in the upwelling region and its adjacent open ocean. Ocean circulation is calibrated with  $\Delta^{14}\text{C}$  data of the ETSP. The sensitivity of the simulated nitrogen cycle to nutrient and oxygen exchange and ventilation from outside the model domain and to remineralization scales inside an OMZ is analysed. For the entire range of model configurations explored, we find that the fixed-N inventory can be stabilized at non-zero levels in the ETSP OMZ only if the remineralization rate via denitrification is slower than that via aerobic respiration. In our optimum model configuration, lateral oxygen supply into the model domain is required at rates sufficient to oxidize at least about one fifth of the export production in the model domain to prevent anoxia in the deep ocean. Under these conditions, our model is in line with the view of phosphate as the ultimate limiting nutrient for phytoplankton, and implies that for the current notion of nitrogen fixation being favoured in N-deficit waters, the water column of the ETSP could even be a small net source of nitrate.

## 2.1 Introduction

The oceanic fixed nitrogen (fixed-N) budget is an essential control on the potential of the ocean to sequester atmospheric  $\text{CO}_2$  via the marine biological pump. Denitrification is generally recognized as a major loss of fixed nitrogen, whereas the balance of the global oceanic nitrogen budget remains controversial. Accordingly, estimates derived from both field data and model analyses for the global oceanic fixed-N budget range from sources roughly balancing sinks [Gruber and Sarmiento, 1997, Gruber, 2004, Eugster and Gruber, 2012, DeVries et al., 2013] to a rather large net deficit between 140 and 234 Tg N yr<sup>-1</sup> [Codispoti et al., 2001, Galloway et al., 2004, Codispoti, 2007].

One of the main uncertainties in the global marine nitrogen budget is the extent of nitrogen loss via denitrification and anaerobic ammonium oxidation (anammox) in oxygen minimum zones (OMZs), located in tropical coastal upwelling regions. Coastal upwelling zones are often associated with very high primary production. Subsequent decomposition of sinking organic matter leads to high levels of oxygen consumption in subsurface waters. Under conditions of sluggish circulation, oxygen-poor source waters, or lack of exchange with oxygenated surface waters, OMZs can develop, usually at intermediate depths of about 200–700 m [Bethoux, 1989, Capone and Knapp, 2007]. An OMZ is commonly defined as a water body with an  $\text{O}_2$  concentration below 20  $\mu\text{mol L}^{-1}$  [Paulmier and Ruiz-Pino, 2009]. The four major open-ocean OMZs

are in the eastern North Pacific (ENP), the eastern tropical South Pacific (ETSP), the Arabian Sea, and the Bay of Bengal. OMZs currently account for only about 8 % of the global ocean area but observations of intense denitrification and anammox in the OMZs indicate that they could be responsible for 30–50 % of the total fixed-N loss [Gruber and Sarmiento, 1997, Codispoti et al., 2001, Dalsgaard et al., 2005, Paulmier and Ruiz-Pino, 2009].

Canfield [2006] used a simple steady-state box model of a coastal OMZ to show that a positive feedback between  $N_2$  fixation and denitrification could strip the OMZ of all fixed nitrogen when  $N_2$  fixation was permitted to restore the nitrate:phosphate ratio to Redfield proportions in the surface ocean. Observed concentrations of fixed-N (nitrate plus nitrite) in OMZ waters, however, typically range from about 15 to 40  $\mu\text{mol L}^{-1}$  [Codispoti and Richards, 1976, Codispoti and Packard, 1980, Morrison et al., 1998, Voss et al., 2001]. A possible explanation for the relatively high nitrate concentrations even in the suboxic core of open-ocean OMZs could be low levels of nitrogen fixation in the overlying surface waters [Landolfi et al., 2013]. However, recent interpretations of observed fixed-N deficits relative to the Redfield equivalent of phosphorus point to high rates of nitrogen fixation closely related to the upwelling of nitrogen-deficit waters along the South American coast [Deutsch et al., 2007]. Although alternative explanations for these nutrient patterns have been proposed in models [Mills and Arrigo, 2010], direct measurements have confirmed the occurrence of nitrogen fixation in and above the OMZ of the ETSP [Fernandez et al., 2011]. Global biogeochemical models also generally predict substantial rates of  $N_2$  fixation in the nitrate-deficit waters of the upwelling region of the ETSP that, if not compensated for by some ad hoc slow-down of remineralization in suboxic conditions, lead to a complete draw-down of nitrate in the OMZ [Moore and Doney, 2007, Schmittner et al., 2008]. The question of how non-zero nitrate concentrations can be maintained in the OMZ thus still awaits a mechanistic answer.

The computational efficiency of box models makes them suitable for sensitivity analyses requiring thousands of model evaluations. In spite of its simplicity, Canfield's (2006) box model is also able to capture the most important first-order interactions among ocean circulation, nitrogen fixation, denitrification and OMZs. However, Canfield's model was limited in its power to investigate the influence of open ocean nutrients and oxygen conditions on the upwelling region, because its dynamics were restricted to the OMZ, where all biogeochemical tracers were prescribed in the surrounding waters. Based on Canfield's (2006) steady-state formulation with prescribed oxygen and nutrient concentrations at all depths outside the OMZ, we here present a fully prognostic box model of  $\text{NO}_3^-$ ,  $\text{PO}_4^{3-}$  and  $\text{O}_2$  cycles in a coastal upwelling region and an adjacent ocean basin. We employ this model to examine under which conditions the observed situation of an essentially complete drawdown of subsurface oxygen and an incomplete drawdown of nitrate can be reproduced for the case of the ETSP. Sensitivity experiments explore how nutrient exchange and oxygen ventilation from the southern boundary influence the nitrogen budget within the model domain.

By including a prognostic description not only of conditions within the OMZ, but also in the water surrounding it, we aim to represent local as well as spatially separated feedbacks between the relatively small OMZ and the much larger open-ocean basin. The model describes net primary and export production by ordinary and  $N_2$ -fixing phytoplankton, as well as aerobic and anaerobic remineralization. The latter is generally associated with nitrogen loss, commonly attributed to denitrification [Codispoti, 1995]. Anammox has recently been reported as another major pathway for fixed-N removal [Kuypers et al., 2005, Hamersley et al., 2007, Molina and Farás, 2009], but the relative contributions of anammox and denitrification are still a matter of debate [Ward et al., 2009, Bulow et al., 2010]. In our model, we do not explicitly resolve the different inorganic nitrogen species and pragmatically consider all loss of fixed-N via anaerobic remineralization as denitrification. In Canfield's (2006) model, most of the physical model parameters were constrained by observations from suboxic upwelling zones in the Arabian Sea and the eastern tropical North and South Pacific. The physical dynamics of our model system are governed by vertical and horizontal mixing and advection, which are calibrated with  $\Delta^{14}C$  data of the ETSP from GLODAP [Global Ocean Data Analysis Project, Key et al., 2004].

## 2.2 Model description

Figure 2.1 shows the model structure, consisting of five boxes representing an upwelling region and an adjacent ocean basin. The U box represents the upper upwelling region. The UM box is the underlying OMZ, where suboxia is expected to develop. The S box represents the surface ocean away from the upwelling zone. Below the S box sits the I box, which represents water of intermediate depth and exchanges water with UM. D is the deep box, which represents water deeper than 500 m. When the UM, I or D boxes become suboxic, denitrification (Denif) will ensue to remineralize the exported organic matter, causing a loss of nitrate (the only form of fixed inorganic nitrogen in our model).

Prognostic tracers represent  $NO_3^-$ ,  $PO_4^{3-}$ ,  $O_2$ ,  $^{14}C$  and the biomass of ordinary and  $N_2$ -fixing phytoplankton, respectively (Table 2.1). The rate of concentration change of a tracer,  $X_i$ , in box  $i$  is composed of physical transport,  $Transport(X_i)$ , and a sources-minus-sinks term,  $SMS(X_i)$ , which represents the effects of biotic processes, air-sea gas exchange and, in the case of  $^{14}C$ , radioactive decay on the tracer concentration (Eq. 2.1).

$$\frac{dX_i}{dt} = \frac{Transport(X_i) + SMS(X_i)}{V_i}$$

$$i \in [U, S, UM, I, D], \quad (2.1)$$

where U, S, UM, I and D refer to the model boxes defined above and  $V_i$  to the corresponding



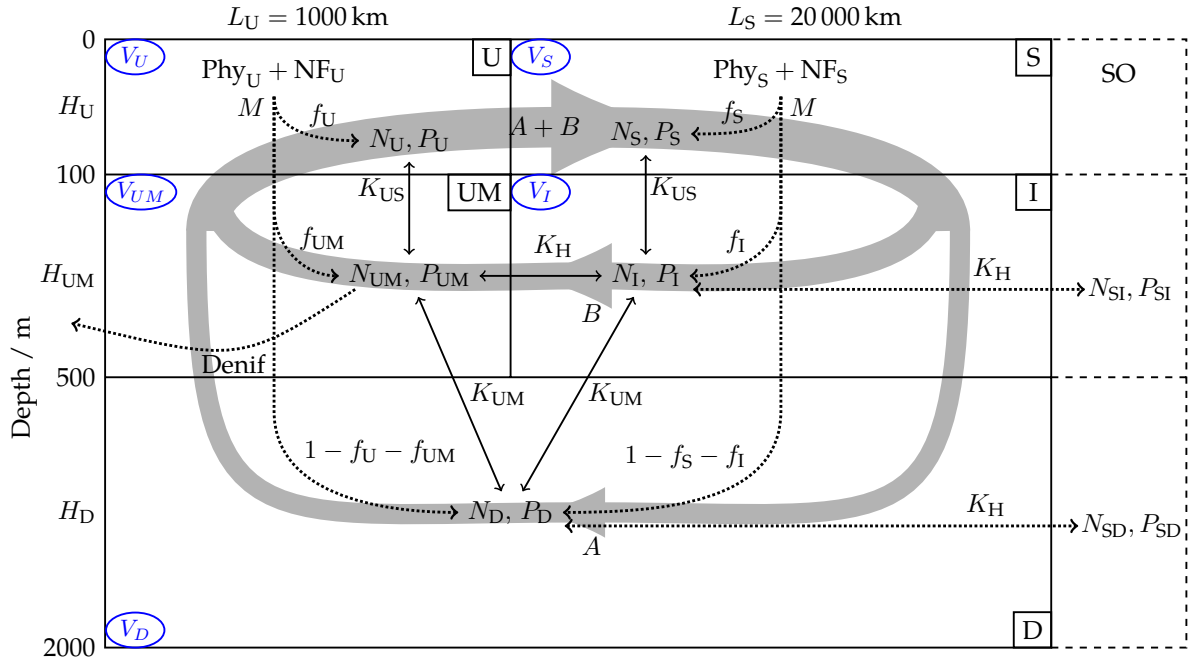


Figure 2.1: Model structure and configurations. The model domain comprises five active boxes representing the top 100 m of an upwelling region (U), the underlying oxygen minimum zone (UM), and an adjacent open-ocean basin divided into a surface (S) and an intermediate-depth box (I). A deep box (D) underlies both the upwelling region and the open ocean. The large-scale circulation is represented by deep ( $A$ ) and shallow ( $B$ ) convection (thick grey lines). Mixing between boxes is implemented via mixing coefficients ( $K$ ). Remineralization derived from net primary production by ordinary (Phy) and diazotrophic (NF) phytoplankton in the surface boxes consumes oxygen. Under anoxic conditions remineralization is fuelled by anaerobic remineralization (Denif). The model can be configured to exchange nutrients and oxygen with the southern subtropical ocean (right, denoted as “SO”). See Table 2.3 for symbol definitions and text for details.

volumes (Fig. 2.1).

### 2.2.1 Transport model

Advection represents the large-scale circulation and is indicated in Fig. 2.1 by wide grey lines.  $L_U$  and  $L_S$  are the horizontal scales of the upwelling region and the open ocean, and  $H_U$ ,  $H_{UM}$  and  $H_D$  refer to the thickness of the respective boxes. The volumes  $V_i$  of the boxes are defined in this 2-D model by  $V_i = H_i \times L_i$ .  $K_{US}$ ,  $K_{UM}$  and  $K_H$  are the coefficients of mixing between different boxes.  $A$  and  $B$  represent the deep and shallow large-scale circulation. The upwelling intensity into box U is given by  $A+B$ . The tracer transport equations for the standard configuration are given in Appendix A (Eqs. 2.12–2.16). All parameters are defined in Tables 2.2 and 2.3.

Variables	Units	Description	Equation
$\text{Phy}_i$	$\mu\text{mol N kg}^{-1}$	Ordinary phytoplankton in box $i$	Eq. (2.2)
$\text{NF}_i$	$\mu\text{mol N kg}^{-1}$	Nitrogen fixers in box $i$	Eq. (2.3)
$\text{N}_i$	$\mu\text{mol N kg}^{-1}$	Nitrate concentration in box $i$	Eq. (2.7)
$\text{N}_{\text{avg}}$	$\mu\text{mol N kg}^{-1}$	Average nitrogen concentration	<sup>a</sup>
$\text{P}_i$	$\mu\text{mol P kg}^{-1}$	Phosphate concentration in box $i$	Eq. (2.8)
$\text{P}_{\text{avg}}$	$\mu\text{mol P kg}^{-1}$	Average phosphorus concentration	<sup>b</sup>
$\text{O}_{2i}$	$\mu\text{mol kg}^{-1}$	Oxygen concentration in box $i$	Eq. (2.9)

$$^a \text{N}_{\text{avg}} = \frac{(\text{N}_U + \text{Phy}_U + \text{NF}_U) \cdot V_U + \text{N}_{UM} \cdot V_{UM} + (\text{N}_S + \text{Phy}_S + \text{NF}_S) \cdot V_S + \text{N}_I \cdot V_I + \text{N}_D \cdot V_D}{V_U + V_{UM} + V_S + V_I + V_D}$$

$$^b \text{P}_{\text{avg}} = \frac{\left(\text{P}_U + \frac{\text{Phy}_U + \text{NF}_U}{r_P}\right) \cdot V_U + \text{P}_{UM} \cdot V_{UM} + \left(\text{P}_S + \frac{\text{Phy}_S + \text{NF}_S}{r_P}\right) \cdot V_S + \text{P}_I \cdot V_I + \text{P}_D \cdot V_D}{V_U + V_{UM} + V_S + V_I + V_D}$$

Table 2.1: Model variables.

Configuration	VID			Units
	STD	VD	VIDRD	
Parameter	OB			
	RD	VDRD	OBRD	
$K_{US}$	8.44	3.37	3.41	$\text{m yr}^{-1}$
$K_{UM}$	1.59	0.40	0.58	$\text{m yr}^{-1}$
$K_H$	47 799	50 475	42 938	$\text{m yr}^{-1}$
$A$	7.20	7.30	7.22	$\text{m yr}^{-1}$
$B$	18.01	19.60	23.07	$\text{m yr}^{-1}$
$g_U$	9.87	8.89	9.88	$\text{m yr}^{-1}$
$g_S$	2.94	1.42	1.46	$\text{m yr}^{-1}$

Configurations in bold are the main configurations.

Table 2.2: Parameters of the physical model configurations. Detailed explanations for these parameters are given in Table 2.3.

### 2.2.2 Biogeochemical model

The ecological model is composed of two phytoplankton types, ordinary phytoplankton, Phy, and nitrogen fixers, NF. The SMS terms for phytoplankton are obtained as the difference between net primary production (NPP) and mortality (M):

$$\text{SMS}(\text{Phy}_i) = (\text{NPP}^{\text{Phy}_i} - \text{M}^{\text{Phy}_i}) \cdot V_i \quad i \in [\text{U}, \text{S}] \quad (2.2)$$

$$\text{SMS}(\text{NF}_i) = (\text{NPP}^{\text{NF}_i} - \text{M}^{\text{NF}_i}) \cdot V_i \quad i \in [\text{U}, \text{S}]. \quad (2.3)$$

Growth of ordinary phytoplankton is described by a Liebig-type dependence on the nitrate and phosphate limitation terms (Eq. 2.4),

$$\text{NPP}^{\text{Phy}_i} = \mu \cdot \min \left( \frac{N_i}{N_i + N_h}, \frac{P_i}{P_i + P_h} \right) \cdot \text{Phy}_i \quad i \in [\text{U}, \text{S}] \quad (2.4)$$

$$\text{NPP}^{\text{NF}_i} = \mu_{\text{NF}} \cdot \frac{P_i}{P_i + P_h} \cdot \text{NF}_i \quad i \in [\text{U}, \text{S}] \quad (2.5)$$

where  $i$  is the model box,  $\mu$  and  $\mu_{\text{NF}}$  are the maximum growth rates of  $\text{Phy}_i$  and  $\text{NF}_i$ , respectively.  $N_i$  and  $P_i$  are nitrate and phosphate concentrations, and  $N_h$  and  $P_h$  are half-saturation concentrations for nitrate and phosphate. Considering possible viral lysis, phytoplankton aggregation, or a feedback between zooplankton grazing and phytoplankton concentration, a quadratic mortality term is adopted for both Phy and NF in all model configurations (Eq. 2.6).

$$\text{M}^{\text{Phy}_i} = M_q \cdot \text{Phy}_i^2 \quad \text{M}^{\text{NF}_i} = M_q \cdot \text{NF}_i^2. \quad (2.6)$$

Table 2.3: Model parameters.

Parameter	Description	Units	Value	Range <sup>(reference)</sup>
$r_a$	O <sub>2</sub> -used/NO <sub>3</sub> -produced during organic carbon(OC) oxidation	—	10.6	8.6–10.6 <sup>a</sup>
$r_c$	C/N ratio of OC oxidation	—	6.63	6.63–7.31 <sup>a</sup>
$f_U$	Remineralization ratio in U	—	20%	b
$f_S$	Remineralization ratio in S	—	20%	b
$f_{UM}$	Remineralization ratio in UM	—	70%	b
$f_I$	Remineralization ratio in I	—	70%	b

Table 2.3: (continued)

Parameter	Description	Units	Value	Range <sup>(reference)</sup>
$r_{\text{den}}$	OC/NO <sub>3</sub> <sup>-</sup> in denitrification	—	1.02	1.02 <sup>a</sup>
$r_{\text{p}}$	N/P released in OC oxidation	—	16	15–16 <sup>a</sup>
$\mu$	Maximum growth rate of Phy	yr <sup>-1</sup>	91.5	36.5–1861.5 <sup>c</sup>
$\mu_{\text{NF}}$	Maximum growth rate of NF	yr <sup>-1</sup>	30.5 <sup>e</sup>	65.7–438 <sup>d</sup>
$M_{\text{q}}$	Quadratic mortality	yr <sup>-1</sup> ( $\mu\text{mol N kg}^{-1}$ ) <sup>-1</sup>	18.25	3.65–18.25 <sup>f</sup>
$N_{\text{h}}$	Nitrate half saturation constant	$\mu\text{mol N kg}^{-1}$	0.5	0.5 <sup>g</sup>
$P_{\text{h}}$	Phosphate half saturation constant	$\mu\text{mol P kg}^{-1}$	0.03125	0.03 <sup>h</sup>
$L_x$	Length of box $x$	m	see Fig. 2.1	—
$H_x$	Depth of box $x$	m	see Fig. 2.1	—
$K_H$	Horizontal exchange	m yr <sup>-1</sup>	n	157.68–56765 <sup>a</sup>
$K_{\text{US}}$	Vertical mixing between surface and intermediate depth	m yr <sup>-1</sup>	n	0.79–31.54 <sup>a</sup>
$K_{\text{UM}}$	Vertical mixing between intermediate depth and deep ocean	m yr <sup>-1</sup>	n	0.21–7.88 <sup>a</sup>
A+B	Upwelling rates	m yr <sup>-1</sup>	n	23.7–630.7 <sup>a</sup>
O <sub>2U</sub>	Oxygen concentration in U	$\mu\text{mol kg}^{-1}$	159.54 <sup>i</sup>	—
O <sub>2S</sub>	Oxygen concentration in S	$\mu\text{mol kg}^{-1}$	198.11 <sup>i</sup>	—
$g_{\text{U}}$	Gas exchange coefficient for U	m yr <sup>-1</sup>	l	—
$g_{\text{S}}$	Gas exchange coefficient for S	m yr <sup>-1</sup>	l	—
$\lambda$	Radioactive decay rate for <sup>14</sup> C	yr <sup>-1</sup>	$1.21 \times 10^{-4}$	$1.21 \times 10^{-4}$ j
N <sub>SD</sub>	Southern boundary nitrate concentration at depth of D	$\mu\text{mol N kg}^{-1}$	32.65	k
N <sub>SI</sub>	Southern boundary nitrate concentration at depth of I	$\mu\text{mol N kg}^{-1}$	10.93	k

Table 2.3: (continued)

Parameter	Description	Units	Value	Range <sup>(reference)</sup>
P <sub>SD</sub>	Southern boundary phosphate concentration at depth of D	$\mu\text{mol N kg}^{-1}$	2.30	k
P <sub>SI</sub>	Southern boundary phosphate concentration at depth of I	$\mu\text{mol N kg}^{-1}$	0.84	k
O <sub>2SD</sub>	Southern boundary oxygen concentration at depth of D	$\mu\text{mol N kg}^{-1}$	181.37	k
O <sub>2SI</sub>	Southern boundary oxygen concentration at depth of I	$\mu\text{mol N kg}^{-1}$	217.98	k

<sup>a</sup>Ranges for  $r_a$ ,  $r_c$ ,  $r_{\text{den}}$ ,  $K_{\text{US}}$ ,  $K_{\text{UM}}$ ,  $K_{\text{H}}$ ,  $A$  and  $B$  are the same as in Canfield [2006].

<sup>b</sup>The fraction of regeneration above 500 m has been estimated between 92% [Suess, 1980] and 97% [Martin et al., 1987]. According to Canfield [2006], most likely 60-70% of the export production is remineralised in the OMZ. Thus, we define 20% and 70% of export production remineralised in the surface boxes and intermediate boxes respectively.

<sup>c</sup>Furnas [1990]

<sup>d</sup>Robarts and Zohary [1987], [Capone et al., 1997]

<sup>e</sup>Temperature-corrected maximum growth rate of NF [Breitbarth et al., 2007].

<sup>f</sup>Palmer and Totterdell [2001]; [Schmittner et al., 2008]

<sup>g</sup>Eppley et al. [1969]

<sup>h</sup>McAllister et al. [1964]

<sup>i</sup>Average 0-100m O<sub>2</sub> concentrations of the corresponding areas from World Ocean Atlas (2009)

<sup>j</sup>Libby [1956]

<sup>k</sup>Averages of the corresponding areas from World Ocean Atlas (2009)

<sup>l</sup>These parameter values are defined in Table 2.2.

Both Phy and NF require phosphate, whereas nitrate is required in addition to phosphate only by Phy, and NF can fix N<sub>2</sub> as long as PO<sub>4</sub><sup>3-</sup> is available (Eqs. 2.4, 2.5). While the ability to utilize organic P has been proposed as an advantage of diazotrophs [Houlton et al., 2008, Ye et al., 2012], ordinary phytoplankton can also use DOP [e.g. Chu, 1946, Cotner, Jr. and Wetzel, 1992] and a clear advantage of diazotrophs over ordinary phytoplankton in the presence of DOP

has never been demonstrated. Thus, we treat all available P to phytoplankton operationally as  $\text{PO}_4^{3-}$  and assume that all organic phosphate is remineralized to  $\text{PO}_4^{3-}$  directly.  $\text{N}_2$  fixers are given a lower maximum growth rate ( $\mu_{\text{NF}}$ , Eq. 2.5), which is 1/3 of the maximum growth rate of ordinary phytoplankton ( $\mu$ ), to account for the high cost of nitrogen fixation [LaRoche and Breitbarth, 2005]. The dependence of NF on iron is not explicitly considered in this model [Mills et al., 2004]. Both types use nitrogen and phosphorus in the Redfield ratio of 16 : 1 [Redfield et al., 1963].

Sensitivity experiments are also performed with a configuration where nitrogen fixers preferentially use nitrate when available and cover only the residual nitrogen demand via  $\text{N}_2$  fixation, denoted as facultative  $\text{N}_2$ -fixation, but overall results are found to be virtually unchanged (Appendix B).

Dead phytoplankton is immediately remineralized in the surface layer and underlying boxes according to the respective remineralization fraction  $f_i$  of box  $i$ . Remineralization occurs preferentially via aerobic respiration, with anaerobic remineralization and the associated nitrogen loss setting in only when all  $\text{O}_2$  has been consumed by aerobic respiration. Phytoplankton growth and aerobic remineralization together with denitrification and, once all inorganic nitrogen is consumed, remineralization via sulfate reduction define the SMS terms of the nitrogen and phosphorus cycles:

$$\text{SMS}(\text{N}_i) = -\text{NPP}^{\text{Phy}_i} \cdot V_i + \text{Rem}_i^{\text{N}} - \text{Denif}_i \quad (2.7)$$

$$\text{SMS}(\text{P}_i) = -\frac{(\text{NPP}^{\text{Phy}_i} + \text{NPP}^{\text{NF}_i}) \cdot V_i}{r_p} + \text{Rem}_i^{\text{P}} \quad (2.8)$$

where the remineralization (Rem) and denitrification (Denif) terms are defined in Appendix C.

$\text{O}_2$  is set constant in the surface ocean boxes U and S, which are in immediate contact with the atmosphere, but varies as a function of transport and respiration in the UM, I and D boxes. Thus, the aerobic respiration terms ( $\text{Res}_i$ ) are only needed in the interior boxes UM, I and D:

$$\text{SMS}(\text{O}_{2i}) = -\text{Res}_i \quad i \in [\text{UM}, \text{I}, \text{D}] \quad (2.9)$$

where Res is defined in Appendix C (Eq. 2.30). Note that aerobic respiration will, in general, also take place in anoxic model boxes, where it utilizes the  $\text{O}_2$  transported from adjacent boxes, before anaerobic respiration starts. All model runs are initialized with  $\text{O}_2$ ,  $\text{NO}_3^-$  and  $\text{PO}_4^{3-}$  annual data from the World Ocean Atlas 2009 [Garcia et al., 2010a,b], averaged over the regions indicated in Table 2.4.

Box	Lat(S)	Lon(W)	Depth(m)	Data <sup>a</sup>
U	5–15	80–90	0–100	–72.39
UM	5–15	80–90	100–500	–93.28
S	0–20	90–190	0–100	–62.21
I	0–20	90–190	100–500	–81.02
D	0–20	80–190	500–2000	–160.30
SI <sup>b</sup>	20–40	90–190	100–500	–71.02
SD <sup>b</sup>	20–40	80–190	500–2000	–134.4

<sup>a</sup>GLODAP “natural  $^{14}\text{C}$ ” data averaged over the respective regions.

<sup>b</sup>SI and SD represent the southern boundary outside the I and D boxes, respectively.

Table 2.4:  $\Delta^{14}\text{C}$  (in ‰) data from GLODAP used for calibration of the model physical parameters.

### 2.2.3 Model calibration

In the present work, the prescribed transport among the different boxes in terms of advection and mixing was calibrated by fitting the modelled  $\Delta^{14}\text{C}$  distribution to the GLODAP “pre-bomb”  $\Delta^{14}\text{C}$  distribution of the eastern tropical South Pacific (Key et al., 2004; Table 2.4). The  $^{13}\text{C}$  fractionation-corrected ratio of  $^{14}\text{C}/^{12}\text{C}$ ,  $\Delta^{14}\text{C}$ , is commonly used in ocean modelling to evaluate and calibrate model physics [Toggweiler et al., 1989, Shaffer and Sarmiento, 1995] because it tends to cancel the effect of the biotic downward transport of  $^{14}\text{C}$  with the rain of organic particles produced by marine organisms. For practical reasons, we employ the arbitrary scale introduced by Shaffer and Sarmiento [1995] to represent  $\Delta^{14}\text{C}$ , and define  $\widehat{\Delta^{14}\text{C}}$ .  $\widehat{\Delta^{14}\text{C}}$  can be converted to  $\Delta^{14}\text{C}$  units as follows:

$$\Delta^{14}\text{C} = 1000(\widehat{\Delta^{14}\text{C}} - 1)\text{‰} \quad (2.10)$$

The  $^{14}\text{C}$  dynamics in the model includes input from the atmosphere at the sea surface and radioactive decay with decay rate  $\lambda = 1.21 \times 10^{-4} \text{ yr}^{-1}$  within the model domain. The SMS term for  $\widehat{\Delta^{14}\text{C}}$  is thus given by:

$$\text{SMS}(\widehat{\Delta^{14}\text{C}}_i) = \left( \frac{g_i(\widehat{\Delta^{14}\text{C}}_a - \widehat{\Delta^{14}\text{C}}_i)}{H_i} - \lambda \widehat{\Delta^{14}\text{C}}_i \right) \cdot V_i \quad i \in [\text{U}, \text{UM}, \text{S}, \text{I}, \text{D}] \quad (2.11)$$

where  $\widehat{\Delta^{14}\text{C}}_i$  is the value of  $\widehat{\Delta^{14}\text{C}}$  in box  $i$ ,  $\widehat{\Delta^{14}\text{C}}_a$  is the atmospheric  $\widehat{\Delta^{14}\text{C}}$ , and  $g_i$  is the gas exchange rate. For  $i \in [\text{UM}, \text{I}, \text{D}]$ ,  $g_i = 0$ . We assume that the  $\Delta^{14}\text{C}$  of the preindustrial atmosphere,  $\Delta^{14}\text{C}_a$ , is 0 ‰. Therefore,  $\widehat{\Delta^{14}\text{C}}_a$  is 1. Model configurations with an open southern boundary also include  $^{14}\text{C}$  exchange with the adjacent subtropical ocean. We also investigate how sensitive our main conclusions are to the uncertainty in the  $\Delta^{14}\text{C}$  data. Table 2.4 shows the  $\Delta^{14}\text{C}$  values used for the calibrations of the different model configurations. Transport and SMS terms for  $^{14}\text{C}$  constitute a system of 5 linear equations with 7 parameters, including 5 transport parameters,  $A$ ,  $B$ ,  $K_{\text{US}}$ ,  $K_{\text{UM}}$ ,  $K_{\text{H}}$ , and 2 air–sea  $^{14}\text{C}$  exchange coefficients,  $g_{\text{U}}$  and  $g_{\text{S}}$ . After setting  $g_{\text{U}}$  and  $g_{\text{S}}$ , the 5 equations can be inverted to solve for the transport parameters. A detailed description of the inversion scheme is provided in Appendix D.

## 2.2.4 Model configurations

The above transport and biogeochemical formulations define the standard (STD) configuration, where the transport and biogeochemical models are applied without exchange with the southern boundary (“SO” in Fig. 2.1). Fixed fractions  $f_i$  of net primary production in U and S are remineralized in the U, UM, S and I boxes, with the remainder being remineralized in D (Fig. 2.1). Nutrient regeneration is assumed to be instantaneous.

In order to investigate the relationships between the different biotic and physical processes and the nitrogen cycle in an OMZ, we introduce seven additional model configurations. The main differences to the STD configuration are shown in Table 2.5.

(1) In the Reduced-denitrification (RD) configuration, denitrification in the OMZ is reduced to 1/5 of the rate of the STD configuration following the procedure applied in the global biogeochemical circulation model by Schmittner et al. [2008]. When oxygen is exhausted in the OMZ, denitrification within the UM box will be responsible for only 1/5 of the remaining organic matter remineralization and the remainder will be remineralized in the D box (Eqs. 2.27 and 2.28). (2) The Ventilation-D (VD) configuration modifies the STD configuration in that the southern boundary of the model domain is partially opened to allow ventilation of  $\text{O}_2$  and  $^{14}\text{C}$  (but not  $\text{NO}_3^-$  and  $\text{PO}_4^{3-}$ ) to the D box from the southern subtropical ocean (Eq. 2.18). The circulation parameters were recalibrated to account for ventilation of  $^{14}\text{C}$  from the south. Ventilation is simulated by applying lateral mixing with the subtropical oligotrophic ocean tracer reservoirs with prescribed  $^{14}\text{C}$  and  $\text{O}_2$  concentrations inferred from observations. All other biogeochemical processes within the model domain are kept the same as in the STD configuration. (3) In the Ventilation-D + Reduced-denitrification (VDRD) configuration, the reduced denitrification rate is applied in the VD configuration. (4) In the Ventilation-ID (VID) configuration, the partially open southern boundary is extended to allow ventilation of  $\text{O}_2$  and  $^{14}\text{C}$  also into the I box from the subtropical ocean (Eq. 2.17). Again, circulation parameters are recalibrated against observed  $\Delta^{14}\text{C}$  taking into account the  $^{14}\text{C}$  exchange between the subtropical



ocean and the I and D boxes. (5) In the Ventilation-ID + Reduced-denitrification (VIDRD) configuration, the reduced denitrification rate is also applied in the VID configuration. (6) In the Open-boundary (OB) configuration, nutrient mixing is added to the VID configuration to allow for the exchange of nutrients between the I and D boxes and the subtropical ocean. (7) In the Open-boundary + Reduced-denitrification (OBRD) configuration, the reduced-denitrification rate is again added to the OB configuration. The physical transports of  $\text{NO}_3^-$  and  $\text{PO}_4^{3-}$  for the I and D boxes in the last six configurations are represented by Eqs. (2.17) and (2.18) in Appendix A.

### 2.2.5 Sensitivity experiments

Two sensitivity experiments are performed for each of the VID and OB configurations to explore the possibilities for preventing  $\text{NO}_3^-$  depletion in the OMZ: (a) different reduced remineralization rates ( $f_{\text{UM}}$ ) and (b) facultative  $\text{N}_2$ -fixation (see Appendix E).

For the OBRD configuration, three sensitivity experiments are performed to investigate our model sensitivity to variable physical transports and biogeochemical tracer concentrations: (1) the mixing rate with the southern boundary,  $K_H$ , is reduced for individual tracers (nutrients, oxygen) or combinations thereof from full rates to zero. (2) Simulations are repeated with individual circulation parameters varied by  $\pm 50\%$  to explore the sensitivity with respect to the circulation parameters of the box model. (3) The sensitivity of  $\text{NO}_3^-$  and  $\text{O}_2$  concentrations in the OMZ to different physical parameters derived from variations of the  $\Delta^{14}\text{C}$  data and  $\text{O}_2$  concentrations in the U-box is also examined.

The literature ranges in Table 2.3 provide only a rough guide for the biogeochemical parameters. The sensitivity of  $r_p$ ,  $\mu$ ,  $\mu_{\text{NF}}$ ,  $N_h$  and  $P_h$  is tested in the OBRD configuration by changing each of them according to the literature range. The effects of changing the remineralization fractions  $f_U$ ,  $f_{\text{UM}}$ ,  $f_S$ , and  $f_I$  are examined by redistributing remineralization between the U and UM, and S and I boxes. These sensitivity experiments will be discussed in Sect. 3.3 below.

## 2.3 Results

The physical circulation parameters calibrated with  $\Delta^{14}\text{C}$  are given in Table 2.2. The resulting circulation is comparable with our expectations for the upwelling region of the eastern tropical South Pacific. The parameters of the biogeochemical model are shown in Table 2.3.

Configuration	Abbreviation	Reduced denitrification rate	O <sub>2</sub> , <sup>14</sup> C ventilation of D	O <sub>2</sub> , <sup>14</sup> C ventilation of I	NO <sub>3</sub> <sup>-</sup> , PO <sub>4</sub> <sup>3-</sup> exchange of D and I
<b>Standard</b>	<b>STD</b>				
<b>Reduced-denitrification</b>	<b>RD</b>	+			
Ventilation-D	VD		+		
Ventilation-D+Reduced-denitrification	VDRD	+	+		
Ventilation-ID	VID		+	+	
<b>Ventilation-ID+Reduced-denitrification</b>	<b>VIDRD</b>	+	+	+	
Open-boundary	OB		+	+	+
<b>Open-boundary+ Reduced-denitrification</b>	<b>OBRD</b>	+	+	+	+

Table 2.5: Summary of model configurations. “+” means that the modification applies to this configuration. The configurations in bold are the main configurations in the text, while the others are the sensitivity configurations described in Appendix E. STD is defined in Sects. 2.2 and 2.3; in RD, a reduced denitrification rate is applied; VD indicates that the southern boundary of the model domain is partially opened to allow ventilation of O<sub>2</sub> and <sup>14</sup>C (but not NO<sub>3</sub><sup>-</sup> and PO<sub>4</sub><sup>3-</sup>) to the D box; VDRD is the configuration when a reduced denitrification rate is applied in VD; VID differs from VD only in that the partially open southern boundary is extended to allow ventilation of O<sub>2</sub> and <sup>14</sup>C also into the I box; VIDRD is the configuration when a reduced denitrification rate is applied in VID; in OB, nutrient (NO<sub>3</sub><sup>-</sup> and PO<sub>4</sub><sup>3-</sup>) mixing is added to VID; OBRD is the configuration in which the reduced denitrification rate is added to OB.

### 2.3.1 Biogeochemical tracer concentrations

Nutrient and oxygen concentrations in the upwelling region (boxes U and UM) are influenced by physical exchange with regions outside the upwelling zone (boxes I and D) and subsequent remineralization of exported organic matter. The results for biogeochemical tracer concentrations of the STD, RD, VIDRD and OBRD configurations are shown in Fig. 2.2 and will be used to develop our main conclusions, whereas those of the VD, VDRD, VID and OB configurations are also included in Fig. 2.3 and described in Appendix E.

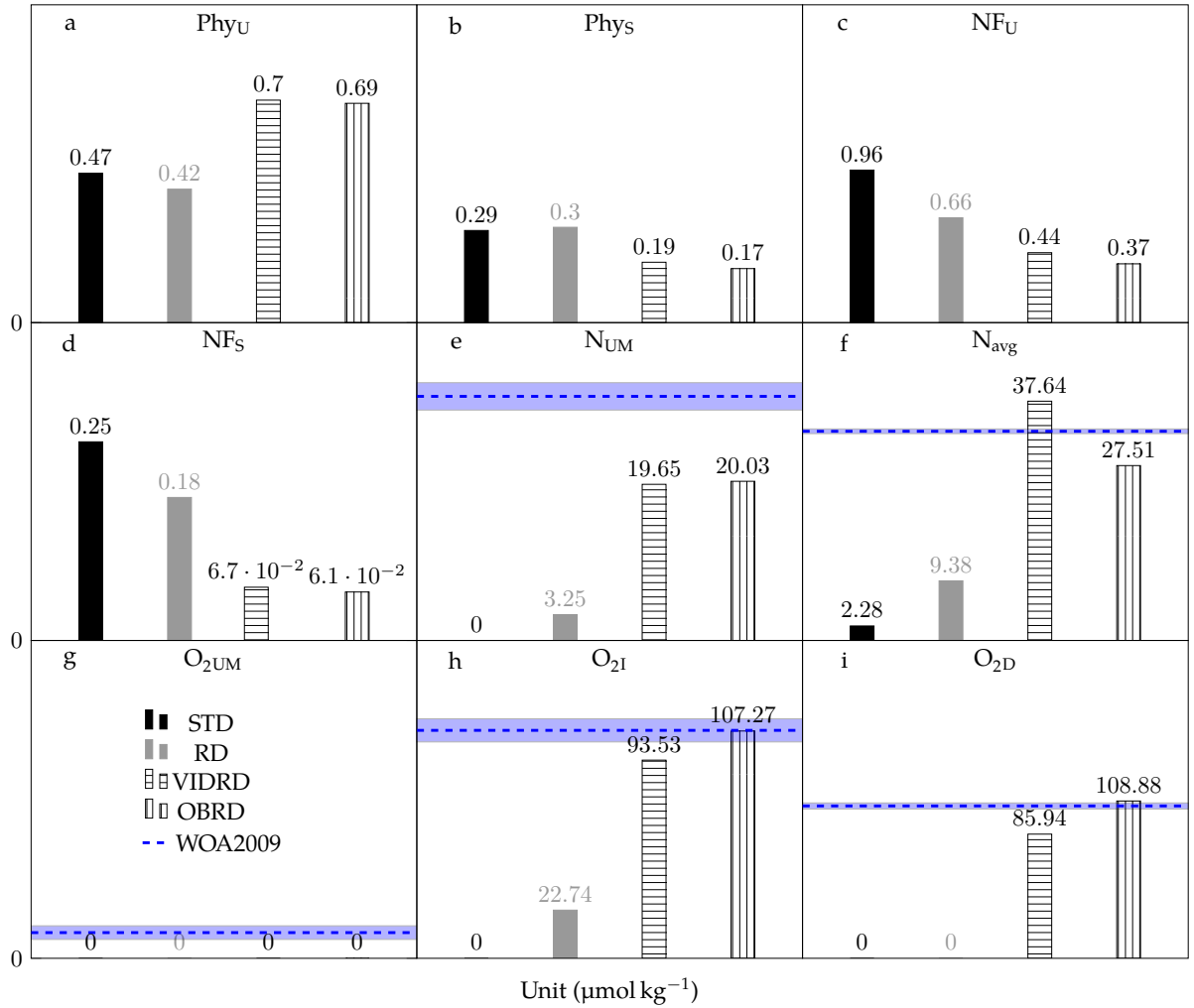


Figure 2.2: Simulated steady-state phytoplankton, nutrient and oxygen concentrations for the main model configurations defined in Tables 2.5 and 2.2. Each panel uses a linear scale of the  $y$  axis starting at zero. Dashed blue lines represent the averages of the WOA2009 nitrate and oxygen data for the corresponding boxes, and the light blue shadings refer to the 95 % confidence intervals; however, there are no data for Phy<sub>U</sub>, Phys, NF<sub>U</sub> and NF<sub>S</sub>.

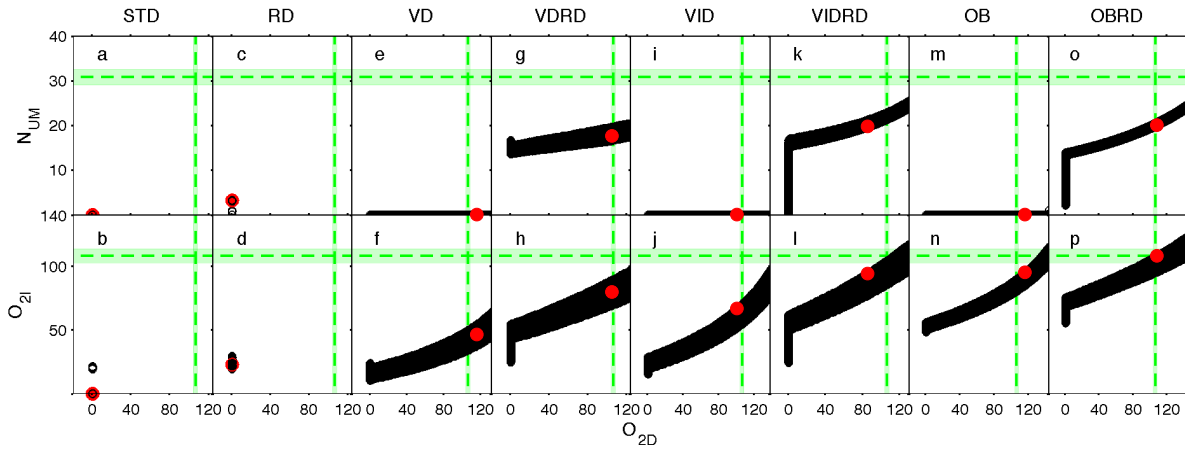


Figure 2.3:  $\text{NO}_3^-$  concentration in the OMZ and  $\text{O}_2$  concentration in the I box for all combinations of  $g_U$  and  $g_S$  resulting in all transport parameters being inside the literature range as given in Table 2.3. The  $x$  axis is the  $\text{O}_2$  concentration in the D box. The red dot in each panel is the selected suite of physical transport parameters which fit the biogeochemical data best in each model configuration. The horizontal green dashed lines represent the averages of the WOA2009 data for  $N_{UM}$  and  $O_{2I}$ , and the vertical green dashed lines denote the averages of the WOA2009 data for  $O_{2D}$ . The light green shadings show the 95 % confidence intervals of the WOA2009 data.

In the STD configuration,  $\text{NO}_3^-$  levels in the UM box are quickly exhausted by denitrification in the UM box (black bars in Fig. 2.2), where decomposition of exported organic matter consumes all  $\text{O}_2$  entering the UM box via advection and diffusion, and all  $\text{NO}_3^-$  entering via physical transport and remineralization of exported organic matter.  $\text{O}_2$  is also depleted in the D box, owing to weak ventilation. This results in over 90 % of the initial nitrate being lost from the model domain by denitrification once the model has reached steady state. The large nitrate deficit with respect to phosphate in the upwelling waters results in nitrogen fixers becoming dominant in the U box despite having a lower maximum growth rate than ordinary phytoplankton (black bars in Fig. 2.2a, c).

In the RD configuration, complete  $\text{NO}_3^-$  exhaustion in the UM box is prevented (grey bars in Fig. 2.2). There is some observational evidence for the efficiency of decomposition of organic matter driven by denitrification in some high-productivity areas being lower than for respiratory processes in the presence of sufficient dissolved oxygen [Liu and Kaplan, 1984, Devol and Hartnett, 2001, Van Mooy et al., 2002]. In our model, this mechanism helps prevent  $\text{NO}_3^-$  depletion in the OMZ because less nitrate is consumed by denitrification during the passage of the particles through the OMZ. However, the  $\text{NO}_3^-$  concentration in the UM box is only  $3.25 \mu\text{mol kg}^{-1}$ , far less than the literature range of about 15 to  $40 \mu\text{mol L}^{-1}$ . The additional organic matter now passing through the OMZ must be remineralized in the D box, which still becomes anoxic in this configuration. Even though the RD configuration fails to simulate the realistic biogeochemical tracer concentrations in the OMZ and its adjacent ocean, it reveals a

possible mechanism for preventing  $\text{NO}_3^-$  depletion in the OMZ.

While a reduced remineralization rate under suboxic conditions appears to be a prerequisite for maintaining non-zero nitrate in the OMZ (Fig. 2.3), we find that ventilation of  $\text{O}_2$  from the lateral subtropical ocean into the model domain is essential for preventing anoxic conditions in the D box. But even ventilation of  $\text{O}_2$  into both I and D boxes fails to prevent  $\text{NO}_3^-$  depletion in the UM box in the VID configuration (Fig. 2.3i). After reducing the denitrification rate in the OMZ, which is the configuration denoted as VIDRD,  $\text{NO}_3^-$  depletion is prevented (bars with horizontal lines in Fig. 2.2). Compared to the RD configuration,  $\text{NO}_3^-$  concentration in the UM box is in a realistic range ( $15\text{--}40 \mu\text{mol kg}^{-1}$ ) for the OMZ, and  $\text{O}_2$  concentrations in the I and D boxes are also closer to the WOA2009 data, but still about 15.0 and 22.7% lower than the observations (blue dashed lines in Fig. 2.2).

In the OBRD configuration, nutrients are exchanged in addition to  $\text{O}_2$  between the subtropical ocean and the I and D boxes, together with reducing the denitrification rate under suboxic conditions.  $\text{N}_{\text{UM}}$  depletion is prevented and  $\text{N}_{\text{UM}}$  approaches a concentration of  $20 \mu\text{mol kg}^{-1}$  at steady state (bars with horizontal lines in Fig. 2.2). The UM box is suboxic and the  $\text{O}_2$  concentrations in the I and D boxes agree very well with the WOA2009 data (blue dashed lines in Fig. 2.2). Overall, the tracer distributions simulated by the OBRD configuration agree best with the observations.

In the sensitivity configuration OB, where only nutrients are exchanged in addition to  $\text{O}_2$  between the subtropical ocean and the I and D boxes, nitrate is still exhausted in the OMZ (Fig. 2.3m).

### 2.3.2 Biogeochemical fluxes

For the biogeochemical fluxes, we focus on the STD, RD, VIDRD and OBRD configurations (configurations in bold in Table 2.5), since they show most clearly the mechanisms responsible for preventing  $\text{NO}_3^-$  depletion in the OMZ (Fig. 2.4). Denitrification rates are much higher in the STD than in the RD configuration (black and grey bars in Fig. 2.4, respectively), even though the physical parameters are the same in the two configurations, leading to nitrate exhaustion in the UM box of the STD but not in the RD configuration. In both VIDRD and OBRD configurations,  $\text{NO}_3^-$  depletion is prevented (bars with horizontal and vertical lines in Fig. 2.2e). Denitrification is weaker in the OBRD than in the VIDRD configuration (bars with horizontal lines in Fig. 2.4), even though the physical parameters are the same in both, keeping more  $\text{NO}_3^-$  in the UM-box in the OBRD configuration. Aerobic respiration in the UM box continues even when the UM box is anoxic. It consumes all  $\text{O}_2$  entering the UM box via the various physical transport processes and thereby oxidizes about 8.7, 64, 85 and 92% of the organic matter remineralized (denitrification + aerobic respiration) in the UM box in all four configurations, STD,

RD, VIDRD and OBRD, respectively. Aerobic respiration is increased in the OBRD configuration compared with the VIDRD configuration (92 % vs. 85 %) at the expense of denitrification.

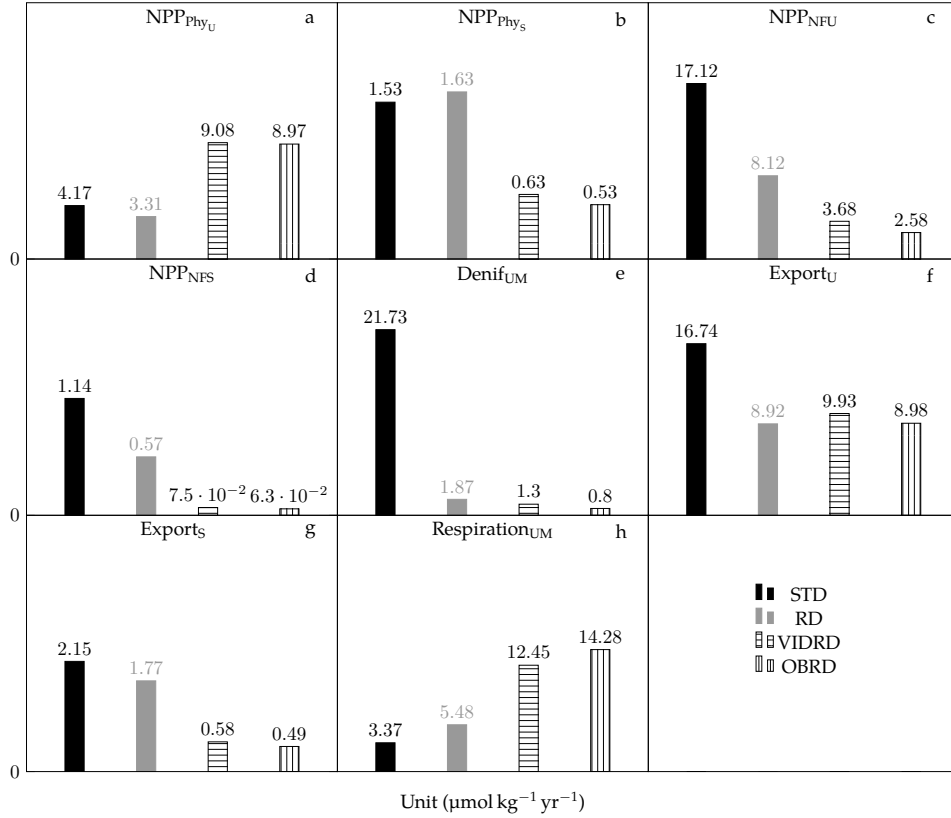


Figure 2.4: Simulated steady-state biogeochemical fluxes for the main model configurations defined in Tables 2.5 and 2.2.  $NPP_{PhyU}$ ,  $NPP_{PhyS}$ ,  $NPP_{NFU}$  and  $NPP_{NFS}$  represent the net primary production rates of  $Phy_U$ ,  $Phy_S$ ,  $NF_U$  and  $NF_S$ , respectively.  $Denif_{UM}$  indicates the nitrogen loss rate by anaerobic remineralization in the UM box.  $Export_U$  and  $Export_S$  represent the export production rate out of U and S.  $Respiration_{UM}$  represents aerobic respiration rate in the UM box. Note that all panels are in units of nitrogen except (h), which is in units of  $O_2$ . Each panel uses a different linear scale for the y axis starting at zero.

Compared with the VIDRD configuration, total  $PO_4^{3-}$  in the model domain decreases by about 28 % in the OBRD configuration at steady state, and the net  $PO_4^{3-}$  flux out of the I box and the corresponding net flux of  $PO_4^{3-}$  from the subtropical ocean into the D box causes a 9.6 % decrease in organic matter exported into the UM box. This explains why  $NO_3^-$  concentrations in the OMZ and  $O_2$  concentrations in the I and D boxes are slightly higher (bars with vertical lines in Fig. 2.2). Restricting, in a hypothetical sensitivity experiment, exchange with the subtropical oligotrophic ocean to  $NO_3^-$  and  $O_2$  does not result in higher  $O_2$  concentrations in the I and D boxes, because  $PO_4^{3-}$  is conserved and export production does not change substantially. Restricting, on the other hand, exchange to  $PO_4^{3-}$  and  $O_2$  depresses production and organic matter decomposition by almost the same amount as in the model run that exchanges  $NO_3^-$ ,  $PO_4^{3-}$  and  $O_2$ , and succeeds in driving  $O_{2I}$  and  $O_{2D}$  concentrations closer to the observations

(not shown). Thus, opening the model boundary to  $\text{PO}_4^{3-}$  and  $\text{O}_2$  helps to improve the model results with respect to those of the VIDRD configuration that do not exchange  $\text{PO}_4^{3-}$  across the southern boundary.

### 2.3.3 Model sensitivity

In sensitivity experiments of the OBRD configurations, designed to elucidate the importance of the influence of the subtropical ocean on the model domain, we reduce the mixing rates ( $K_H$ ) of  $\text{O}_2$ ,  $\text{NO}_3^-$  and  $\text{PO}_4^{3-}$  across the southern boundary in different combinations (two of them are shown in Fig. 2.5). Total net primary production (NPP) is the same when varying the mixing rate of either only  $\text{O}_2$  or  $\text{O}_2$  together with  $\text{NO}_3^-$ , as long as the  $\text{PO}_4^{3-}$  exchange rate with the subtropical ocean remains unchanged (dotted lines in Fig. 2.5b). The reason is that changes in nitrate exchange with the subtropical ocean are compensated for by changes in denitrification and nitrogen fixation. This implies that NPP is, at steady state, determined only by phosphate availability. Once the total NPP of NF and Phy is determined by  $\text{PO}_4^{3-}$  supply, aerobic respiration will increase with increasing  $\text{O}_2$  supply, while anaerobic remineralization will decrease (solid and dashed lines in Fig. 2.5c). Comparing simulations that vary the mixing rate of  $\text{O}_2$  together with that of  $\text{PO}_4^{3-}$ , with simulations that vary the mixing rate of  $\text{O}_2$  together with that of  $\text{NO}_3^-$  and  $\text{PO}_4^{3-}$  (dotted lines in Fig. 2.5g) reveals that the combined NPP of Phy and NF decreases with decreasing  $\text{PO}_4^{3-}$  inventory at low mixing rates and increases with increasing  $\text{PO}_4^{3-}$  inventory at high mixing rates, irrespective of  $\text{NO}_3^-$  mixing rate.

It turns out that  $\text{NO}_3^-$  depletion in the OMZ is prevented in the four sensitivity experiments of Fig. 2.5 no matter how much  $\text{O}_2$  is supplied from the subtropical ocean into the model domain. But anoxia in the D box can only be prevented if the external  $\text{O}_2$  supply can oxidize more than about 20 % export production in the model domain (solid lines in Fig. 2.5e, j). This ratio can vary according to the size of S box. It increases for a smaller S box, because a smaller S box leads to less downward  $\text{O}_2$  supply from the surface box into the lower model domain and more lateral ventilation is needed, and vice versa (not shown). These experiments also reveal an interesting link among the  $\text{O}_2$  supply from the subtropical ocean, the nitrate concentration in the OMZ, and the  $\text{NO}_3^-$  flux between the model domain and surrounding ocean. As soon as the  $\text{O}_2$  supply from the subtropical ocean accounts for about 17.5 % of the respiratory  $\text{O}_2$  consumption, the model domain turns into a small net source of  $\text{NO}_3^-$  (dashed lines in the first row of Fig. 2.5). Once anoxia is prevented in the D box, the nitrate concentration in the OMZ (UM box) increases quite strongly from about  $15 \mu\text{mol kg}^{-1}$  to reach about  $20 \mu\text{mol kg}^{-1}$  at full mixing strength.

The conclusion that the model domain is a small pelagic net  $\text{NO}_3^-$  source in the OBRD configuration does not change when individual physical transport parameters vary by up to  $\pm 50\%$  (Fig. 2.6). Varying biogeochemical parameters also does not affect this conclusion. The finding

that the model domain is a net  $\text{NO}_3^-$  source is also tenable for all suites of physical transport parameters in the literature range for which the D-box is oxic (Fig. 2.7).

The oxygen concentrations in the I and D boxes increase in sensitivity model runs with physical parameters calibrated from increased  $^{14}\text{C}$  concentrations (lower water mass age, figure not shown).  $\text{NO}_3^-$  depletion is prevented in the OMZ in the OBRD configuration within the range of about  $\pm 10\%$  around the default ventilation intensity (Fig. 2.8a). The UM box remains anoxic when  $\Delta^{14}\text{C}$  is varied within this range, which indicates that this mechanism for preventing  $\text{NO}_3^-$  depletion in the OMZ may be robust against modest changes in the regional ocean circulation patterns. Meanwhile,  $\text{NO}_3^-$  and  $\text{O}_2$  in the UM box are quite insensitive to the  $\text{O}_2$  concentration in the U-box for this configuration (Fig. 2.8b).

Varying biogeochemical parameters affects individual model predictions but not our main conclusions. The strongest effects are those of varying the N:P ratio  $r_p$  and the remineralization fractions ( $f_U$ ,  $f_{UM}$ ,  $f_S$  and  $f_I$ ) (Fig. 2.6). Lowering  $r_p$  to 12 increases  $N_{UM}$  by about 35 %, but cannot change the strength of our model domain as a net  $\text{NO}_3^-$  source. Increasing  $r_p$  to 20 decreases  $N_{UM}$  by about 18 %, but triples the strength of our model domain as a  $\text{NO}_3^-$  source. However, observations indicate that  $r_p$  for the ETSP is more likely to be higher than lower compared to the Redfield N:P ratio of 16 [Franz et al., 2012]. Increasing the maximum growth rate of NF,  $\mu_{NF}$ , to  $1/2\mu$ , the maximum growth rate of Phy, results in higher  $N_{UM}$  concentrations and our model domain being a larger  $\text{NO}_3^-$  source. Intuitively, decreasing  $\mu_{NF}$  to  $1/4\mu$  results in lower  $N_{UM}$  concentrations and our model domain becoming a smaller  $\text{NO}_3^-$  source. Varying the  $\text{NO}_3^-$  half saturation constant,  $N_h$ , results in virtually unchanged results.  $N_{UM}$  increases when changing remineralization fractions in the intermediate boxes ( $f_{UM}$  and  $f_I$ ) from 70 to 50 and 30 %, respectively, effectively lowering export production via lowering the export ratio. Nevertheless, the qualitative behaviour of the model remains the same in these sensitivity experiments.

## 2.4 Discussion and conclusions

Nitrogen is often considered to be the primary limiting nutrient in marine upwelling regions with OMZs [Cline and Richards, 1972, Codispoti and Christensen, 1985, Morrison et al., 1998, Voss et al., 2001], where denitrification rates can be high and are generally thought to cause a major loss of  $\text{NO}_3^-$  from the world's oceans. The nitrogen deficit is commonly assumed to



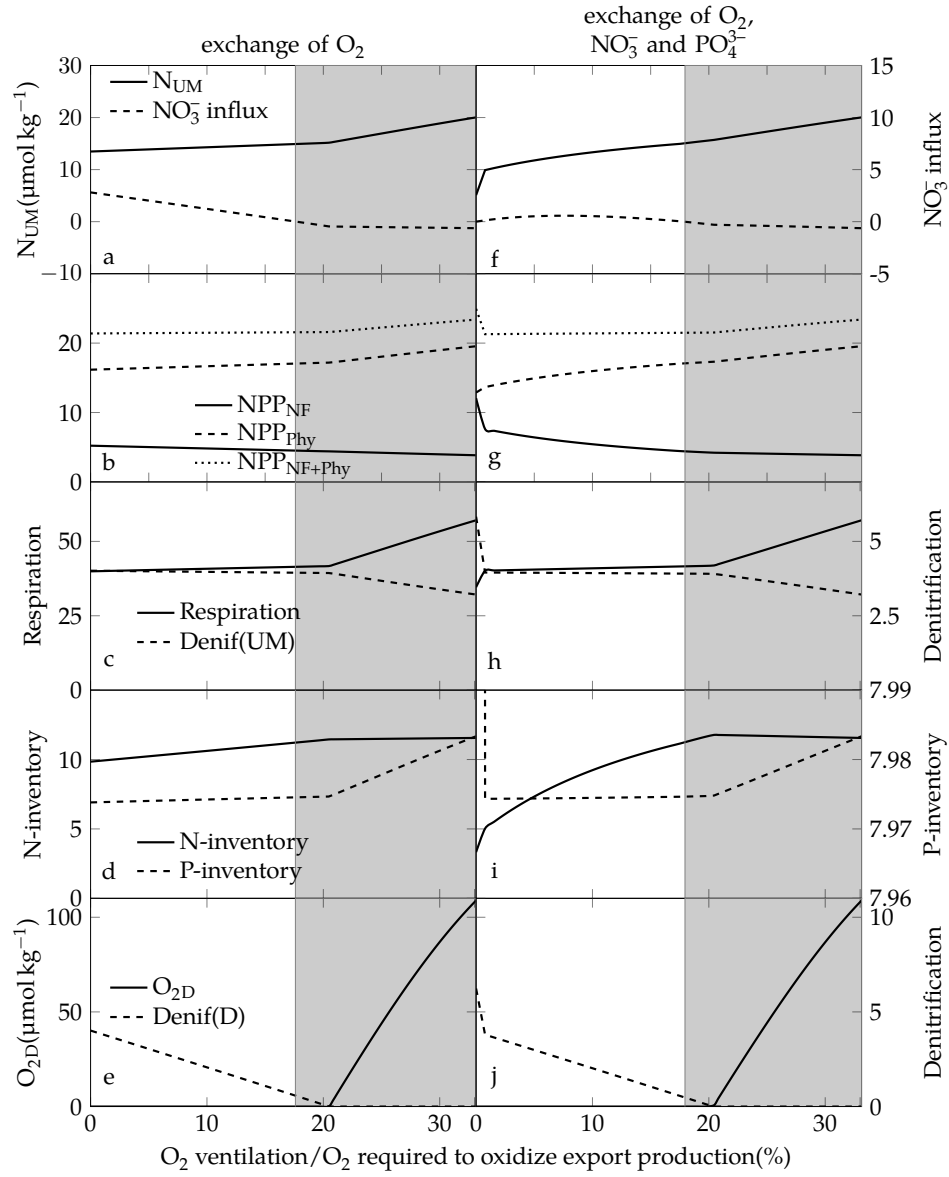


Figure 2.5: Dependence of biogeochemical processes on the exchange of  $O_2$ ,  $NO_3^-$  and  $PO_4^{3-}$  with the subtropical ocean through the southern boundaries of the I and D boxes. The  $x$  axes indicate the contribution of  $O_2$  supplied from the subtropical ocean relative to that required to oxidize all export production from the surface ocean (boxes U and S). (a–e) only  $O_2$  exchanged through the southern boundaries is reduced; (f–j) exchange of  $O_2$ ,  $NO_3^-$  and  $PO_4^{3-}$  is reduced.  $N_{UM}$  is  $NO_3^-$  concentration in the UM box and  $NO_3^-$  influx is the  $NO_3^-$  flux through the southern boundary (positive into model domain).  $NPP_{Phy}$ ,  $NPP_{NF}$  and  $NPP_{NF+Phy}$  are net primary production by ordinary phytoplankton, nitrogen fixers, and the sum of both in the surface ocean. Respiration and Denif(UM) represent  $O_2$  consumption by aerobic remineralization and  $NO_3^-$  removal by anaerobic remineralization, respectively, in the UM box. N-inventory and P-inventory are the total nitrogen and phosphorus inventories in the model domain, including all organic and inorganic species.  $O_{2D}$  and Denif(D) represent  $O_2$  concentration and  $NO_3^-$  removal by anaerobic remineralization in the D box. Units of all variables are  $10^{11} \mu\text{mol yr}^{-1} \text{m}^{-1}$  except for  $N_{UM}$  and  $O_{2D}$ , which are given in  $\mu\text{mol kg}^{-1}$ , and N-inventory and P-inventory, which are  $10^{11} \mu\text{mol m}^{-1}$ . The shaded area denotes the parameter range for which the model domain is a net source of  $NO_3^-$ .

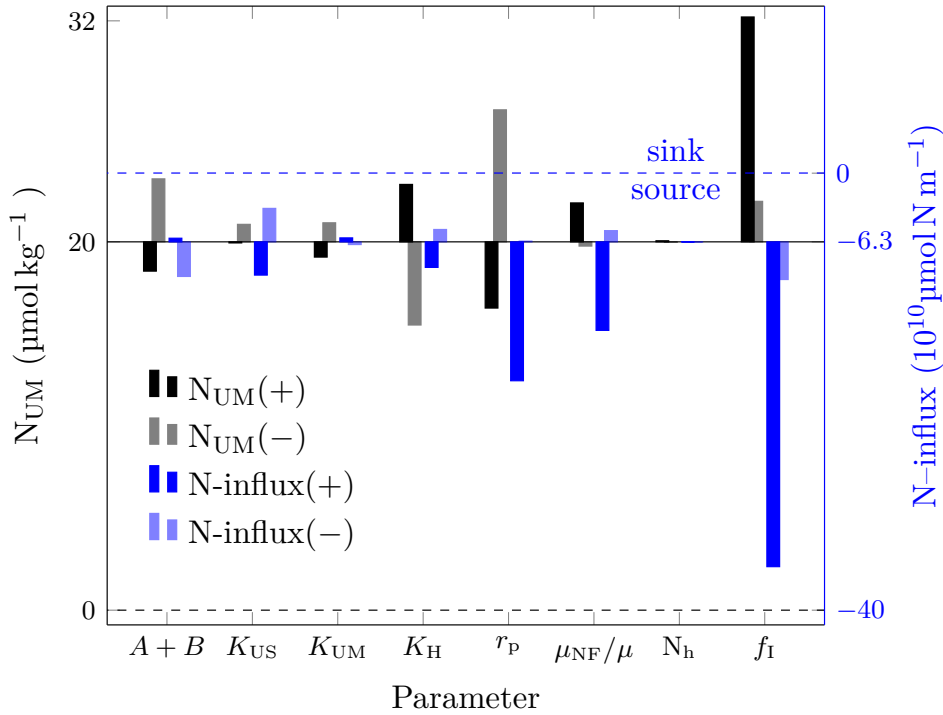


Figure 2.6: Sensitivity of  $\text{NO}_3^-$  concentration in the OMZ ( $N_{UM}$ ) and the net  $\text{NO}_3^-$  flux out of the model domain to variations of the individual parameters describing ocean transport and biogeochemical processes (see Tables 2.2, 2.3 and Fig. 2.1 for a description of the parameters). Black and blue bars represent changes in  $N_{UM}$  and  $N\text{-influx}$ , respectively. “+” and “-” indicate the response to increased and decreased parameters. Physical circulation parameters are varied by  $\pm 50\%$ .  $r_p$  is varied between 12 and 20.  $\mu_{NF}/\mu$  is varied between  $1/4$  and  $1/2$ .  $N_h$  varies between  $0.3$  and  $0.9 \mu\text{mol kg}^{-1}$ . For  $f_I$ , “+” indicates  $f_U = f_S = 60\%$  and  $f_{UM} = f_I = 30\%$ , and “-” means  $40\%$  and  $50\%$ , respectively.

stimulate  $\text{N}_2$  fixation, both in observational estimates of  $\text{N}_2$  fixation [Deutsch et al., 2007, Monteiro et al., 2010] and in current descriptions of  $\text{N}_2$  fixation in biogeochemical models [Moore and Doney, 2007, Schmittner et al., 2008]. However, if  $\text{N}_2$  fixation is tightly linked to nitrogen loss processes, denitrification of organic matter derived from  $\text{N}_2$  fixation can consume more nitrogen than was fixed via  $\text{N}_2$  fixation and thereby lead to a vicious cycle of runaway nitrogen loss [Landolfi et al., 2013]. This has been found to lead to total depletion of  $\text{NO}_3^-$  in the OMZ of box models [Canfield, 2006] and global biogeochemical circulation models [Moore and Doney, 2007, Schmittner et al., 2008]. For our fully prognostic 5-box model, we have identified the mechanisms capable of arresting the runaway nitrogen loss that can result from a close coupling of nitrogen fixation and denitrification [Landolfi et al., 2013] and maintaining realistic non-zero  $\text{NO}_3^-$  concentrations in open-ocean OMZs of the ETSP: reduced  $\text{NO}_3^-$  consumption by denitrification owing to slower remineralization under suboxic compared to oxic conditions, coupled with lateral ventilation of oxygen and, to some extent, nutrients with the subtropics.

In this work, the model configuration with reduced denitrification rate, and lateral ventilation and nutrient exchange (OBRD configuration) performs best with respect to the observations,

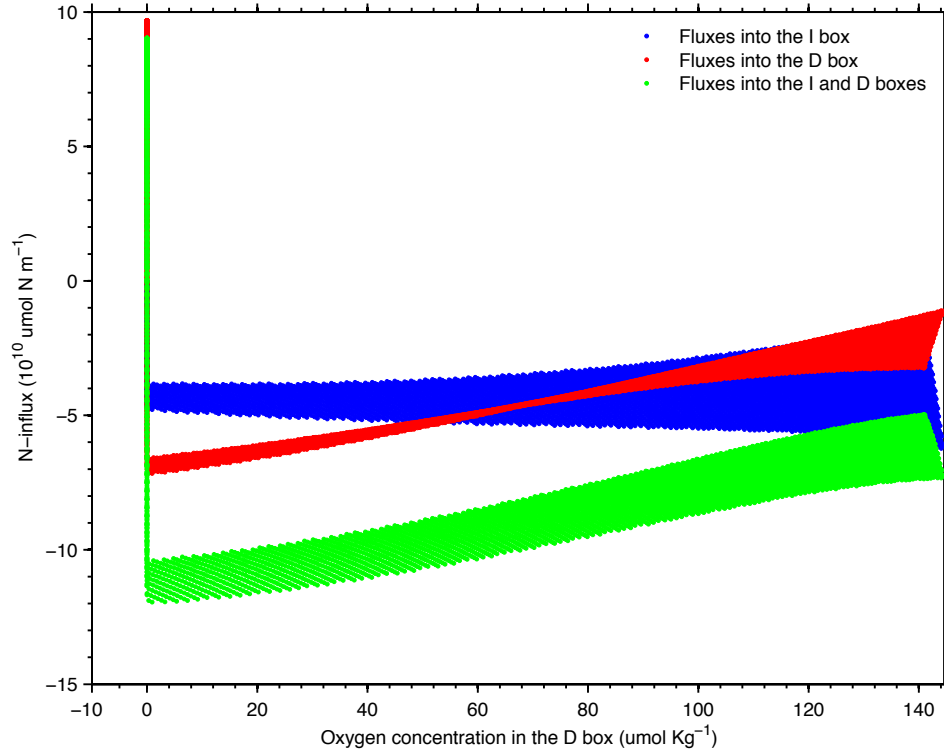


Figure 2.7: Lateral  $\text{NO}_3^-$  input into the model domain of the OBRD configuration as a function of the oxygen concentration in the D box. “Fluxes into the I box” represents lateral  $\text{NO}_3^-$  input into the I box; “fluxes into the D box” represents lateral  $\text{NO}_3^-$  input into the D box; “fluxes into the I and D boxes” is the sum of the above two processes. Note that only the I and D boxes can exchange  $\text{O}_2$  or nutrients with the region outside of the model domain. In this figure, negative values indicate that the model is a  $\text{NO}_3^-$  source, and positive values that the model is a  $\text{NO}_3^-$  sink instead.

whereas reduced rates of denitrification alone lead to unrealistic  $\text{O}_2$  depletion in the deep waters, and reduced rates of denitrification combined with only  $\text{O}_2$  ventilation in the D, or both I and D boxes still underestimate the WOA2009 data for  $\text{O}_2$  in the I and D boxes. By allowing for exchange of  $\text{O}_2$  and nutrients with the subtropical ocean in the OBRD configuration, the delicate balance between sufficient  $\text{O}_2$  supply required for maintaining high levels of aerobic respiration and sufficient  $\text{PO}_4^{3-}$  loss through the open boundary appears fulfilled. Compared with other box models [Shaffer and Sarmiento, 1995, Tyrrell, 1999, Deutsch et al., 2004, Canfield, 2006, Mills and Arrigo, 2010, Eugster and Gruber, 2012, DeVries et al., 2012], the box model we use here explicitly employs both a reduced-denitrification rate and an open boundary condition, which seems to be a prerequisite for the ability to simulate realistic nitrate conditions in the OMZ and oxygen concentrations in the adjacent ocean.

The  $\text{NO}_3^-$  loss by denitrification in the OMZ of the OBRD configuration is compared with that of other model-based and observational estimates in Table 2.6. Our simulated denitrification is consistent with the results of Somes et al. [2010] and DeVries et al. [2013] for the ETSP, lower

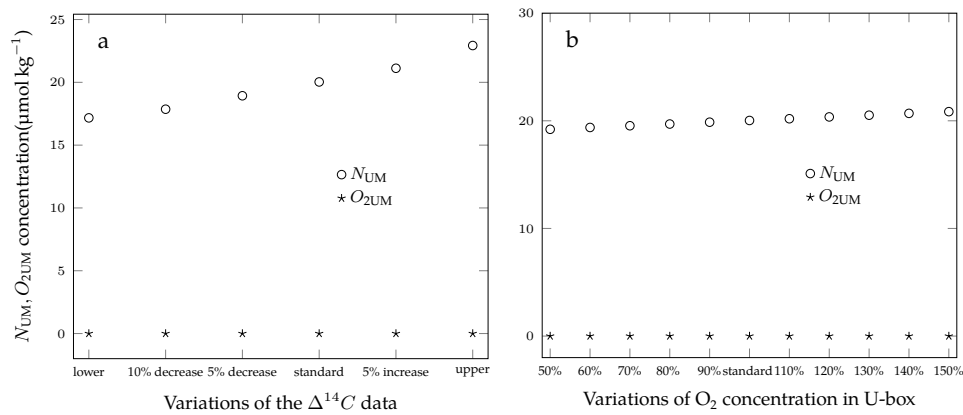


Figure 2.8:  $\text{NO}_3^-$  and  $\text{O}_2$  concentrations in the OBRD configuration for different physical parameters derived from variations of the  $\Delta^{14}\text{C}$  data **(a)** and  $\text{O}_2$  concentration in the U box **(b)**. **(a)** Decrease and increase mean that  $\Delta^{14}\text{C}$  values in all boxes are reduced or increased simultaneously. **(b)** Values of the  $x$  axis denote the variations of  $\text{O}_2$  concentration in the U box relative to the standard. The standard run in each figure is the OBRD configuration with physical parameters defined in Table 2.2.

than the estimate of Bianchi et al. [2012] and Kalvelage et al. [2013], but higher than that of Mills and Arrigo [2010]. However, the estimated denitrification by Bianchi et al. [2012] represents that of the entire South Pacific but not only the ETSP. The Kalvelage et al. (2013) model has much higher fixed-N influx into the OMZ via physical transport than our model, which could compensate for their more intense  $\text{NO}_3^-$  loss by denitrification.

Data/models	Denitrification ( $\text{Tg N yr}^{-1}$ )
OBRD configuration	5.0 <sup>c</sup>
Kalvelage et al. [2013] <sup>a</sup>	10.2 <sup>c</sup>
T. DeVries et al. (2013, pers. comm.) <sup>b</sup>	$7.0 \pm 2.0$ <sup>c</sup>
Bianchi et al. [2012] <sup>a</sup>	17.6 <sup>d</sup>
C. Somes et al. (2010, pers. comm.) <sup>b</sup>	5.8 <sup>c</sup>
Mills and Arrigo [2010] <sup>b</sup>	1.9 <sup>c,*</sup>

Table 2.6: Denitrification comparison with model-based and observational estimates. <sup>a</sup> Observational estimate; <sup>b</sup> model results; <sup>c</sup> ETSP; <sup>d</sup> entire South Pacific; \* OMZ value extrapolated to the UM box of our model.

Phosphate turns out to be the ultimate limiting nutrient in our model (Fig. 2.5), and hence determines the nitrogen content of the OMZ. This strong control of the N cycle by phosphate is similar to the findings of previous models [e.g. Lenton and Watson, 2000, Canfield, 2006],

where the occurrence and extent of oceanic anoxia was also tightly linked to phosphate supply. These studies explicitly assumed that  $\text{N}_2$  fixation acts to restore surface-oceanic dissolved inorganic N:P ratio towards the Redfield ratio. By contrast, we do not prescribe the effect of nitrogen fixation on surface inorganic nutrients, and the finding of phosphate as the ultimate limiting nutrient is obtained regardless of whether NF responds directly to the N:P ratio in the surface ocean or whether NF is inhibited by the presence of  $\text{NO}_3^-$  (see Appendix B).

Even though our model indicates only a weak dependence of simulated  $\text{NO}_3^-$  concentrations in the OMZ on the lateral  $\text{O}_2$  supply from the subtropical ocean,  $\text{O}_2$  depletion in the adjacent ocean can be prevented only when the  $\text{O}_2$  supply from the subtropical ocean accounts for more than 20 % of the  $\text{O}_2$  required to oxidize export production from the surface ocean of our model domain (U and S boxes). This value only varies between about 17 and 23 %, when the size of the S box varies by  $\pm 50$  %, indicating that our above conclusion is rather insensitive to our choice of model domain. The  $\text{O}_2$  supply from the subtropical ocean might be also linked to the balance of the regional fixed-N cycle. In the most realistic (OBRD) configuration, the balance of water-column denitrification and nitrogen fixation is tightly linked to the nitrate concentration in the OMZ: if the model domain acts as a sink for  $\text{NO}_3^-$ , nitrate concentrations in the OMZ do not exceed about  $15 \mu\text{mol kg}^{-1}$ , i.e. the lower limit of current observations, 15 to  $40 \mu\text{mol kg}^{-1}$  [Codispoti and Richards, 1976, Codispoti and Packard, 1980, Morrison et al., 1998, Voss et al., 2001]. Nitrate concentrations close to those commonly found in OMZs are predicted only when nitrogen fixation exceeds water-column denitrification and the ocean basin containing the OMZ becomes a net source of  $\text{NO}_3^-$  (Fig. 2.5a, f).

There are no specific data or model results focusing on the water-column nitrogen balance of the ETSP. Our simulations using a parameterization where nitrogen fixation is inhibited by the presence of nitrate yield very similar results, only with slightly reduced nitrogen fixation resulting in a somewhat reduced (by  $\leq 15$  %) nitrogen source for strong enough lateral oxygen supply. Ganachaud and Wunsch [2002] estimated a net northward  $\text{NO}_3^-$  transport of  $270 \pm 170 \text{ kmol s}^{-1}$  ( $119.2 \pm 75.1 \text{ Tg N yr}^{-1}$ ) across  $17^\circ\text{S}$  into the ETSP in a geostrophic inverse box model, which indicates that the ETSP is a net nitrogen sink, but their estimate included benthic denitrification, which is not accounted for in our current analysis. In a model-guided analysis, DeVries et al. [2012] predicted water-column denitrification rates of  $21\text{--}33 \text{ Tg N yr}^{-1}$  by simulating the distribution of observed dissolved  $\text{N}_2$  gas produced by denitrifying bacteria. From an ocean circulation-biogeochemical model-based analysis of nutrient concentrations and transport rates, Deutsch et al. [2007] estimated nitrogen fixation rates in the Pacific Ocean of about  $95 \text{ Tg N yr}^{-1}$ , half of which was speculated to occur in the ETSP. From these estimates, we cannot rule out that the water column of the ETSP is a net source of  $\text{NO}_3^-$ , which would be consistent with our results obtained in the OBRD configuration. More recently, Eugster and Gruber [2012] probabilistically estimated nitrogen fixation and water-column and benthic denitrification separately in their box model, which appears to be consistent with our results as their results also indicate that the water column of the Indo-Pacific is a large fixed-N source.

Based on our model results, we conclude that reduced  $\text{NO}_3^-$  consumption by denitrification owing to slower remineralization under suboxic conditions together with lateral transport is essential to arrest the vicious cycle of runaway fixed-N loss in the OMZ. More research is needed to better constrain the model parameters, in particular the upwelling transport and the difference between the organic matter remineralization rate via aerobic respiration and anaerobic denitrification.

## 2.5 Appendix A

Physical transports of a tracer  $X$  among the boxes U, UM, S, I and D are defined for the STD configuration as follows:

$$\text{Transport}(X_U) = (X_{UM} - X_U)(A + B + K_{US}) \cdot L_U \quad (2.12)$$

$$\begin{aligned} \text{Transport}(X_{UM}) = & [AX_D + BX_I - X_{UM}(A + B) + K_{US}(X_U - X_{UM})] \cdot L_U \\ & + K_{UM}(X_D - X_{UM}) \cdot L_U + K_H(X_I - X_{UM}) \cdot H_{UM} \end{aligned} \quad (2.13)$$

$$\text{Transport}(X_S) = (A + B)(X_U - X_S) \cdot L_U + K_{US}(X_I - X_S) \cdot L_S \quad (2.14)$$

$$\begin{aligned} \text{Transport}(X_I) = & (A + B)(X_S - X_I) \cdot L_U + K_H(X_{UM} - X_I) \cdot H_{UM} \\ & + K_{US}(X_S - X_I) \cdot L_S + K_{UM}(X_D - X_I) \cdot L_S \end{aligned} \quad (2.15)$$

$$\text{Transport}(X_D) = A(X_I - X_D) \cdot L_U + K_{UM}(X_{UM} - X_D) \cdot L_U + K_{UM}(X_I - X_D) \cdot L_S \quad (2.16)$$

For the VD, VID and OB configurations, Eqs. 2.15 and 2.16 are modified to read:

$$\begin{aligned} \text{Transport}(X_I) = & (A + B)(X_S - X_I) \cdot L_U + K_H(X_{UM} - X_I) \cdot H_{UM} \\ & + K_{US}(X_S - X_I) \cdot L_S + K_{UM}(X_D - X_I) \cdot L_S \\ & + K_H(X_{SI} - X_I) \cdot H_{UM} \end{aligned} \quad (2.17)$$

$$\begin{aligned} \text{Transport}(X_D) = & A(X_I - X_D) \cdot L_U + K_{UM}(X_{UM} - X_D) \cdot L_U + K_{UM}(X_I - X_D) \cdot L_S \\ & + K_H(X_{SD} - X_D) \cdot H_D \end{aligned} \quad (2.18)$$

The transport equations (Eqs. 2.12, 2.14) are modified for Phy and NF because Phy and NF are assumed to be capable of regulating their buoyancy and exist only in U and S without being transported vertically between the U and UM or the S and I boxes.

## 2.6 Appendix B

It is well known from laboratory studies that diazotrophic phytoplankton can also utilize nitrate for growth, denoted as facultative  $N_2$ -fixation [e.g. Holl and Montoya, 2005]. Schmittner et al. [2008] introduced a formulation where nitrogen fixers preferentially use nitrate when available and cover only the residual nitrogen demand via  $N_2$  fixation. In order to examine the behaviour of our model when nitrogen fixers (NF) preferentially use nitrate, nitrogen fixation is separated from diazotroph growth, as in Schmittner et al. [2008]:

$$\text{Nitrogen-fixation}_i = \left(1 - \frac{N}{N + N_h}\right) \cdot \text{NPP}^{\text{NF}_i} \quad i \in [\text{U}, \text{S}] \quad (2.19)$$

and Eq. (2.7) was modified to accommodate the additional nitrate uptake by NF as follows:

$$\begin{aligned} \text{SMS}(N_i) &= (-\text{NPP}^{\text{Phy}_i} - (\text{NPP}^{\text{NF}_i} - \text{Nitrogen-fixation}_i)) \cdot V_i + \text{Rem}_i^N - \text{Denif}_i \\ i &\in [\text{U}, \text{UM}, \text{S}, \text{I}, \text{D}] \end{aligned} \quad (2.20)$$

After incorporating Eqs. 2.19 and 2.20 in our all configurations described in Sect. 2.2, they behave very similarly compared to Fig. 2.2, except that  $\text{Phy}_U$  and  $\text{Phy}_S$  concentrations are somewhat lower because nitrogen fixers take up  $\text{NO}_3^-$  as well. Thus, we conclude that our results are robust with respect to assumptions about facultative  $N_2$ -fixation by diazotrophs.

## 2.7 Appendix C

The nitrogen cycle in this model includes nitrogen fixation, denitrification, inorganic nitrogen regeneration by remineralization, physical transport of  $\text{NO}_3^-$ , and transport of Phy and NF between the U and S boxes. The physical transport process of  $\text{NO}_3^-$  is the same as for other tracers.  $\text{Rem}_i^N$  represents nitrogen remineralization in box  $i$  with fixed fractions  $f_i$  of the net

primary production in the overlaying surface boxes.

$$\text{Rem}_U^N = f_U M(\text{Phy}_U + \text{NF}_U) \cdot H_U \cdot L_U \quad (2.21)$$

$$\text{Rem}_{UM}^N = f_{UM} M(\text{Phy}_U + \text{NF}_U) \cdot H_U \cdot L_U \quad (2.22)$$

$$\text{Rem}_S^N = f_S M(\text{Phy}_S + \text{NF}_S) \cdot H_U \cdot L_S \quad (2.23)$$

$$\text{Rem}_I^N = f_I M(\text{Phy}_S + \text{NF}_S) \cdot H_U \cdot L_S \quad (2.24)$$

$$\text{Rem}_D^N = M[(1 - f_U - f_{UM})(\text{Phy}_U + \text{NF}_U) \cdot L_U + (1 - f_S - f_I)(\text{Phy}_S + \text{NF}_S) \cdot L_S] \cdot H_U. \quad (2.25)$$

Denitrification (Denif) is the nitrogen loss to  $\text{N}_2$  when UM, I and D are anoxic. It is defined as:

$$\text{Denif}_i = \max(\text{Rem}_i^N - \frac{\text{Res}_i}{r_a}, 0) \cdot \frac{r_c}{r_{\text{den}}} \quad i \in [\text{UM}, \text{I}, \text{D}] \quad (2.26)$$

with the aerobic respiration term  $\text{Res}_i$  given below (Eq. 2.30). In the RD, VDRD, VIDRD and OBRD configurations, the equations for denitrification in the  $i$  box and remineralization in D are given by:

$$\text{Denif}_i = \frac{1}{5} \cdot \max(\text{Rem}_i^N - \frac{\text{Res}_i}{r_a}, 0) \cdot \frac{r_c}{r_{\text{den}}} \quad i \in [\text{UM}, \text{I}] \quad (2.27)$$

$$\begin{aligned} \text{Rem}_D^N &= M[(1 - f_U - f_{UM})(\text{Phy}_U + \text{NF}_U) \cdot L_U + (1 - f_S - f_I)(\text{Phy}_S + \text{NF}_S) \cdot L_S] \cdot H_U \\ &+ 4 \cdot \frac{\text{Denif}_i \cdot r_{\text{den}}}{r_c} \quad i \in [\text{UM}, \text{I}]. \end{aligned} \quad (2.28)$$

All the organic matter is composed according to the Redfield ratio, i.e. C:N:P = 106 : 16 : 1.  $\text{Rem}_i^P$  represents phosphate remineralization in box  $i$ , which is obtained by dividing  $\text{Rem}_i^N$  by the Redfield N:P ratio,  $r_P$ :

$$\text{Rem}_i^P = \frac{\text{Rem}_i^N}{r_P} \quad i \in [\text{U}, \text{S}, \text{UM}, \text{I}, \text{D}]. \quad (2.29)$$

Respiration ( $\text{Res}_i$ ) is considered in the intermediate and deep ocean. In the presence of sufficient oxygen or when oceanic circulation supplies enough  $\text{O}_2$ , organic matter will be preferentially oxidized by respiration. We use a ratio of  $-\text{O}_2 : \text{N} = 170 : 16$  during oxic remineralization of organic matter [Anderson and Sarmiento, 1994].

$$\text{Res}_i = \begin{cases} r_a \cdot \text{Rem}_i^N & \text{if } \text{O}_{2i} > 0 \\ \min(\text{Transport}(\text{O}_{2i}), r_a \cdot \text{Rem}_i^N) & \text{if } \text{O}_{2i} = 0 \end{cases} \quad (2.30)$$



$$i \in [\text{UM}, \text{I}, \text{D}].$$

## 2.8 Appendix D

Transport and SMS terms for  $\widehat{^{14}\text{C}}$  constitute a system of 5 linear equations including transport parameters  $A$ ,  $B$ ,  $K_{\text{US}}$ ,  $K_{\text{UM}}$  and  $K_{\text{H}}$ , which are inverted from the  $\Delta^{14}\text{C}$  values with air–sea  $\Delta^{14}\text{C}$  exchange rates  $g_{\text{U}}$  and  $g_{\text{S}}$  for the U and S boxes, respectively as inputs. All possible combinations of values (with a step size of  $0.01 \text{ m yr}^{-1}$ ) for  $g_{\text{U}}$  and  $g_{\text{S}}$  are applied to derive the values for transport parameters  $A$ ,  $B$ ,  $K_{\text{US}}$ ,  $K_{\text{UM}}$  and  $K_{\text{H}}$ .  $g_{\text{U}}$  and  $g_{\text{S}}$  are constrained in a two-step procedure. First, all combinations are determined which result in transport parameters in the literature range in Table 2.3. Finally, the combination giving the most realistic biogeochemical tracer concentrations (closest to observations) is chosen for the experiments in the main text (Fig. 2.3). This approach for determining the physical transport parameters is applied to obtain those in Table 2.2.

## 2.9 Appendix E

The VD, VDRD, VID and OB configurations are sensitivity configurations that we employ to explore the ability of several mechanisms for preventing nitrogen exhaustion in the OMZ. Even though all of them fail in simulating realistic biogeochemical tracer concentrations compared with observations, they reveal insights into the effects of the respective mechanisms.

The VD configuration, which differs from the STD configuration in supplying additional  $\text{O}_2$  from the subtropical ocean into the D box, avoids anoxic deep waters, but fails to prevent nitrate exhaustion in the UM box (Fig. 2.3e, f). The I and D boxes are oxic at steady state, but  $\text{NO}_3^-$  is still exhausted by denitrification in the UM box. Comparing the oxygen concentrations of the RD and VD configurations, we conclude that ventilation helps to prevent oxygen depletion in the D box, but this mechanism alone cannot prevent eventual runaway denitrification in the UM box.

After incorporating reduced denitrification in the VDRD configuration, nitrate depletion is prevented and the I and D boxes are oxic (Fig. 2.3g, h). But the  $\text{O}_2$  concentration in the I box is about 28% lower than that of the WOA2009 data, which indicates that  $\text{O}_2$  supply into the D-box only may not be sufficient to ventilate the model domain.

The VID configuration, which is modified from the VD configuration by including  $\text{O}_2$  ventilation into the I box, fails to prevent  $\text{NO}_3^-$  depletion by denitrification in the OMZ (Fig. 2.3i, j).

In the OB configuration, nutrients are exchanged in addition to  $O_2$  between the subtropical ocean and the I and D boxes. In this configuration, nitrate exhaustion in the OMZ is still not prevented, even though phytoplankton and nitrogen fixer concentrations are lower in the surface U and S boxes (Fig. 2.3m, n).

Two further sensitivity experiments were performed for each of the VID and OB configurations to explore how  $NO_3^-$  depletion in the UM box can be prevented. (1) Decreasing the fraction of export production remineralized in the UM box ( $f_{UM}$ ) from 70 to 56 % makes  $NO_3^-$  persist in the UM box. Together with the 20 % remineralization in the U box, this implies that 76 % of the export production is remineralized in the upper 500 m of the ocean. However, the resulting  $NO_3^-$  concentration in the UM box is far below the literature range of about 15 to 40  $\mu\text{mol L}^{-1}$ . (2) Facultative  $N_2$ -fixation inhibits nitrogen fixation in an environment with high  $NO_3^-$  concentrations, but fails to prevent  $NO_3^-$  depletion in the UM box.

The sensitivity with respect to the biogeochemical parameters (Sect. 2.5) is examined for all configurations. We only discuss the sensitivity of the OBRD configuration in Sect. 3.3, since the sensitivity experiments in all model configurations do not affect our main conclusion that fixed-N inventory can be stabilized at non-zero levels in the ETSP OMZ only if the remineralization rate via denitrification is slower than that via aerobic respiration. The model behaviour is very similar in all sensitivity experiments, with two exceptions. When  $\mu_{NF}$  is raised to  $1/2\mu$  and remineralization fractions in different boxes are rearranged to  $f_U=f_S=60\%$  combined with  $f_{UM}=f_I=30\%$ , the VDRD and VIDRD configurations reach a steady state only in combination with facultative  $N_2$ -fixation.

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## Chapter 3

# Box-modeling of the impacts of atmospheric nitrogen deposition and benthic remineralization on the nitrogen cycle of the eastern tropical South Pacific

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**Abstract** Both atmospheric deposition and benthic remineralization influence the marine nitrogen cycle, and hence ultimately also marine primary production. The biological and biogeochemical relations of the eastern tropical South Pacific (ETSP) to nitrogen deposition, benthic denitrification and phosphate regeneration are analysed in a prognostic box model of the oxygen, nitrogen and phosphorus cycles in the ETSP. In the model, atmospheric nitrogen deposition based on estimates for the years 2000-2009 is offset by half by reduced  $N_2$  fixation, with the other half transported out of the model domain. Both model- and data-based benthic denitrification are found to trigger nitrogen fixation, partly compensating for the  $NO_3^-$  loss. Since phosphate is the ultimate limiting nutrient in the model, enhanced sedimentary phosphate regeneration under suboxic conditions stimulates primary production and subsequent export production and  $NO_3^-$  loss in the oxygen minimum zone (OMZ). A sensitivity analysis of the local response to both atmospheric deposition and benthic remineralization indicates dominant stabilizing feedbacks in the ETSP, which tend to keep a balanced nitrogen inventory, i.e., nitrogen input by atmospheric deposition is counteracted by decreasing nitrogen fixation;  $NO_3^-$  loss via benthic denitrification is partly compensated by increased nitrogen fixation; enhanced nitrogen fixation stimulated by phosphate regeneration is partly removed by the stronger water-column denitrification. Even though the water column in our model domain acts as a  $NO_3^-$  source, the ETSP including benthic denitrification might become a  $NO_3^-$  sink.

### 3.1 Introduction

Marine primary production (PP) by phytoplankton is a key factor controlling the strength of the oceanic biological carbon pump and the amount of  $CO_2$  that is stored in the ocean [Gruber, 2004, Okin et al., 2011]. PP is controlled by light and nutrients, such as nitrogen, phosphorus or iron, necessary for the production of phytoplankton. These nutrients are supplied to the light-lit surface waters by upwelling, turbulent entrainment of subsurface water, riverine inputs, biological nitrogen fixation, atmospheric deposition and benthic remineralisation [Falkowski et al., 1998, Kasai et al., 2002, Duce et al., 2008, Bakun and Weeks, 2008, Moore and Braucher, 2008].

Nitrogen is often the limiting nutrient for phytoplankton in the ocean [Moore et al., 2013]. On the other hand, oceanic nitrogen is thought to adjust, via nitrogen gain and loss processes, to the marine phosphorus inventory on geological time scales, making phosphorus the ultimate limiting nutrient and nitrogen the proximate limiting nutrient [Tyrrell, 1999]. The ocean's nitrogen inventory has a turnover time of a few thousand years, being affected by relatively large interacting nitrogen sinks and sources. The exact mechanisms and timescales of the interactions are not well understood. Estimates of oceanic nitrogen fixation, the main fixed-N source into the ocean, vary from 106 to 330 Tg N yr<sup>-1</sup> based on both in-situ observations and

models [Codispoti et al., 2001, Brandes and Devol, 2002, Gruber and Sarmiento, 2002, Gruber, 2004, Großkopf et al., 2012]. Water-column denitrification and anaerobic ammonium oxidation (anammox) in oxygen minimum zones (OMZs), accounting for  $100\text{--}300\text{ Tg N yr}^{-1}$ , and benthic denitrification, estimated as  $95\text{--}300\text{ Tg N yr}^{-1}$ , mainly determine the oceanic fixed-N sink [Gruber and Sarmiento, 2002, Gruber, 2004, Codispoti, 2007, Bohlen et al., 2012, Eugster and Gruber, 2012, DeVries et al., 2012]. Due to the large uncertainty in the major sources and sinks of the global nitrogen cycle, the balance of the nitrogen inventory in the ocean is still a matter of debate [Gruber, 2004, Codispoti, 2007, DeVries et al., 2012].

Phosphate can be the ultimate limiting nutrient on geological time scales even in regions with fixed nitrogen deficits with respect to the Redfield-equivalent of the phosphate concentration [Tyrrell, 1999, Su et al., 2015, Auguères and Loreau, 2015]. The ocean's phosphorus budget has been suggested to be unbalanced in the modern ocean with sedimentary burial as the major sink exceeding phosphorus sources [Wallmann, 2010]. This condition might be alleviated by benthic phosphate regeneration, which can be enhanced under low-oxygen bottom waters ( $\text{O}_2 < 20\text{ }\mu\text{mol L}^{-1}$ ) [Slomp and Van Cappellen, 2007, Wallmann, 2010]. Input of bioavailable phosphorus into the ocean stimulates primary production, and decomposition of subsequent export production enhances  $\text{O}_2$  consumption in the ocean, in turn increasing the volume of oceanic oxygen-deficit water and the fixed-N loss. Consequently, phosphate regeneration is expected to be enhanced by enlarging OMZs, possibly leading to a positive feedback loop [Van Cappellen and Ingall, 1994, Wallmann, 2003].

OMZs also play an important role in the global marine fixed-N budget as they are responsible for a large fraction of total marine fixed-N loss [Canfield, 2006]. The relative contribution of heterotrophic denitrification and autotrophic anammox to the total oceanic fixed-nitrogen sink remains debated [Lam et al., 2009, Ward et al., 2009]. Even though anammox has been observed to be a major fixed-N loss in eastern tropical South Pacific (ETSP) [Lam et al., 2009, Kalvelage et al., 2013], both denitrification and anammox are ultimately driven by the flux of organic matter into the OMZ [Koeve and Kähler, 2010, Kalvelage et al., 2013]. For simplicity, heterotrophic denitrification is considered the major fixed-N loss process in the present study. Continental shelves and the upper continental slopes are the most important sites for benthic fixed-N loss [Christensen et al., 1987, Devol, 1991]. However, Bohlen et al. [2011] found that the continental shelf and upper continental slope of the ETSP across a section at  $11^\circ\text{S}$  are sites of nitrogen recycling rather than fixed-N loss, because of relatively low rates of denitrification and high rates of  $\text{NH}_4^+$  release from Dissimilatory Nitrate Reduction to Ammonium (DNRA). This illustrates that the  $\text{NH}_4^+$  released from DNRA should be taken into account when the benthic fixed-N sink is estimated.

In the last few decades, a number of model and data based investigations have been carried out on the importance of atmospheric fixed-N input into the ocean for marine biogeochemical cycles [Duce, 1986, Duce et al., 1991, Krishnamurthy et al., 2007, 2010, Okin et al., 2011,

Mouriño-Carballido et al., 2012]. Duce et al. [2008] indicate that anthropogenic nitrogen deposition is rapidly approaching the global oceanic estimates for  $N_2$  fixation, while preindustrial deposition was an order of magnitude lower than  $N_2$  fixation. However, the response of nitrogen-fixation and denitrification to atmospheric nitrogen deposition remains an open question. Atmospheric nitrogen inputs into the global ocean are dominated by inorganic nitrogen from anthropogenic sources [Warneck, 1988, Paerl and Whittall, 1999]. The exact magnitude of organic nitrogen deposition is not clear due to a lack of observations [Cornella et al., 2003, Duce et al., 2008, Zamora et al., 2011]. Thus, the contribution of DON to total nitrogen deposition is still uncertain and the distribution, bioavailability and lifetime are also not clear [Seitzinger and Sanders, 1999, Duarte et al., 2006, Duce et al., 2008]. Therefore, DON deposition is excluded from our analysis.

Various biogeochemical models have addressed the effects and feedbacks between the major sources and sinks in the marine nitrogen cycle [Van Cappellen and Ingall, 1994, Deutsch et al., 2001, 2007, Krishnamurthy et al., 2007, Somes et al., 2013, Landolfi et al., 2013]. However, most of them have explored only a subset of the atmospheric, pelagic and benthic nitrogen sources and sinks. Using a conceptually simple and computationally efficient box model, we here attempt a synthesis considering all essential sources and sinks and their mutual interactions, with the only exception of riverine input, which is excluded from our model analysis because of its unknown and presumably minor role in the ETSP.

## 3.2 Model description

### 3.2.1 Circulation and biogeochemical model

The circulation model is the same as in Su et al. [2015], which is a prognostic 5-box model to explore the interactions among oceanic circulation, nitrogen fixation and water-column denitrification in the OMZ of the ETSP. Briefly, the physical parameters were calibrated to fit the average  $\delta^{14}C$  of each box and biogeochemical parameters are constrained by literature data. All the simulations in this manuscript employ the Open-boundary + Reduced-denitrification (OBRD) configuration of Su et al. [2015], which allows for exchange of deep and intermediate ETSP waters with the southern subtropical ocean ("SO" in Fig. 3.1) and applies reduced remineralisation rates under suboxic conditions. The model domain consists of five boxes representing the water column of an upwelling region and an adjacent ocean basin. The U box represents the upper upwelling region. The UM box is the OMZ below, where suboxia is expected to develop. The S box represents the surface ocean away from the upwelling zone. Below the S box sits the I box, which represents water of intermediate depth and exchanges water with UM. D is the deep box, which represents water deeper than 500 m (model configuration shown in Fig. 3.1).

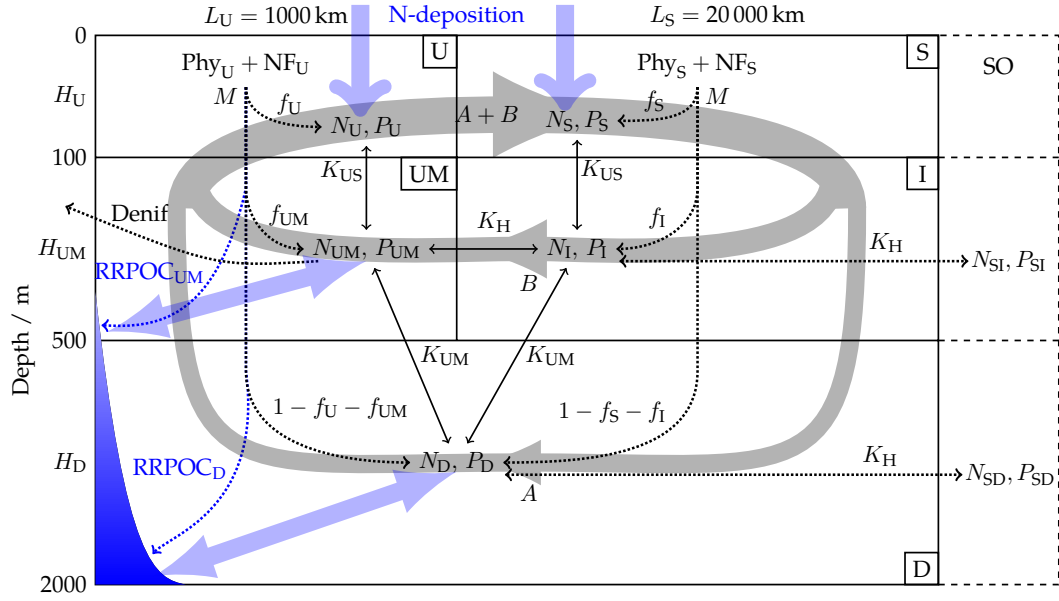


Figure 3.1: Model structure. The model domain comprises five boxes representing the top 100 m of an upwelling region (U), the underlying oxygen minimum zone (UM), and an adjacent open-ocean basin divided into a surface (S) and an intermediate-depth box (I). A deep box (D) underlies both the upwelling region and the open ocean. The large-scale circulation is represented by deep (A) and shallow (B) convection (thick grey lines). Mixing between boxes is implemented via mixing coefficients ( $K$ ). Remineralisation derived from primary production by ordinary (Phy) and diazotrophic (NF) phytoplankton in the surface boxes consumes oxygen. Under anoxic conditions remineralisation is fueled by anaerobic remineralization (Denif). In the configuration employed in this study, the model domain exchanges nutrients and oxygen with the southern subtropical ocean (right, denoted as “SO”). Nitrogen deposition and benthic remineralization are included additionally to represent their influence on the local water-column nutrient concentrations (thick light blue arrows).

There are two phytoplankton types in the biogeochemical model, ordinary phytoplankton, Phy, and nitrogen fixers, NF, as defined in Su et al. [2015]. Both Phy and NF require phosphate, whereas nitrate is required in addition to phosphate only by Phy, and NF can fix  $N_2$  as long as  $PO_4^{3-}$  is available. Dead phytoplankton is immediately remineralized in the surface layer and underlying boxes according to the pre-defined remineralization fractions. Remineralization occurs preferentially via aerobic respiration, with anaerobic denitrification and the associated nitrogen loss setting in only when all  $O_2$  has been consumed by aerobic respiration. When oxygen is exhausted in the OMZ, remineralization is assumed to slow down by a factor of 5, and accordingly denitrification within the UM box is responsible for 1/5 of the remaining organic matter remineralization and the remainder will be remineralized in the D box.

In order to represent the nitrogen and phosphate fluxes across the water–sediment interface, remineralization of particulate organic carbon reaching the sediment (POC rain rate, RRPOC) is included additionally in the UM and D boxes. RRPOC is calculated according to the method

Configuration	Process				
	Model BD	Data BD	Model PR	Data PR	N-DEP
Control					
N-DEP					+
MBD	+				
MPR			+		
DBD		+			
DPR				+	
Synthesis configuration					
MBD+MPR (Syn1)	+		+		
DBD+DPR (Syn2)		+		+	
MBD+MPR+N-DEP (Syn3)	+		+		+
DBD+DPR+N-DEP (Syn4)		+		+	+

Table 3.1: Summary of model configurations. ‘Model BD’ and ‘Data BD’ represent model- and data-based benthic denitrification, respectively. ‘Model PR’ and ‘Data PR’ are model- and data-based benthic phosphate regeneration, respectively. ‘N-DEP’ represents the atmospheric nitrogen input into the surface ocean according to the estimate by Lamarque et al. [2011]. ‘+’ indicates that the process is included.

introduced in Section 3.2.4, and we assume that all the POC is buried in the sediment.

### 3.2.2 Model configurations

The above descriptions define the control configuration. In order to investigate the model sensitivity to atmospheric nitrogen deposition and benthic remineralization, we employ another nine model configurations incorporating either a subset or all of these processes, which are summarized in Table 3.1. Detailed information of all processes is presented in Sections 3.2.3, 3.2.4 and 3.2.5.

### 3.2.3 Atmospheric nitrogen deposition

Years 2000–2009 levels of dry and wet inorganic nitrogen deposition following the RCP 4.5 scenario [Lamarque et al., 2011] are examined in our work. Inferred atmospheric inorganic nitrogen deposition rates are  $0.081$  and  $1.4 \text{ Tg N yr}^{-1}$  ( $73.1$  and  $64.9 \text{ mg N m}^{-2} \text{ yr}^{-1}$ ) for the U and S box, respectively. Note that the circulation remains constant in our model, and only



atmospheric nitrogen deposition fluxes are included as an additional annual nitrogen input into the surface (U and S) boxes.

Duce et al. [1991] estimated atmospheric dissolved inorganic phosphorus (DIP) input into the global ocean, which indicates high N/P (mole/mole) ratios of more than 100 on a global scale [Dentener et al., 2006, Duce et al., 2008, Mahowald et al., 2008], much higher than the average elemental N/P required by phytoplankton in the ocean. Thus, we do not explore the influence of atmospheric phosphorus deposition in our analysis.

### 3.2.4 Benthic denitrification

The empirical transfer function of Bohlen et al. [2012] is applied to predict benthic inorganic nitrogen loss ( $L_{\text{DIN}}$  in  $\mu\text{mol N m}^{-2} \text{d}^{-1}$ ) through benthic denitrification, which can account for the net loss of dissolved inorganic nitrogen (DIN) from the sediment.

$$L_{\text{DIN}} = (0.06 + 0.19 \times 0.99^{(\text{O}_2 - \text{NO}_3^-)_{\text{bw}}}) \times \text{RRPOC} \quad (3.1)$$

where  $\text{NO}_3^-$  and  $\text{O}_2$  are bottom-water nitrate and oxygen concentrations in  $\mu\text{mol kg}^{-1}$ , and the rain rate (RRPOC) is in  $\mu\text{mol C m}^{-2} \text{d}^{-1}$ . Since the bottom-water  $\text{NO}_3^-$  and  $\text{O}_2$  concentrations are well known in the ETSP, the uncertainty in our estimation of benthic denitrification comes mostly from uncertainties in the rain rate, which, in turn, depends on biological production, as a function of phytoplankton biomass and its physiological status. Simulated phytoplankton concentrations in the surface boxes of the model roughly agree with estimates by Behrenfeld et al. [2005] from Aqua-Modis satellite data and the Redfield C:N ratio (U-box:  $1.06 \mu\text{mol N kg}^{-1}$  simulated vs.  $0.68 \mu\text{mol N kg}^{-1}$  from Aqua-Modis; S-Box:  $0.23 \mu\text{mol N kg}^{-1}$  simulated vs.  $0.28 \mu\text{mol N kg}^{-1}$  from Aqua-Modis).

### Model-based estimation of benthic denitrification

Fixed-N losses via benthic denitrification ( $L_{\text{DIN}}$ ) in the UM and D boxes are obtained according to Eq. 3.1, with the respective simulated actual  $\text{NO}_3^-$  and  $\text{O}_2$  concentrations taken as the bottom-water concentrations, and RRPOC is estimated from the export production out of the U and S boxes ( $\text{EP}_\text{U}$  and  $\text{EP}_\text{S}$ ) and the Martin curve (Eq. 3.2) [Martin et al., 1987]:

$$\text{RRPOC} = F \times \left( \frac{z}{100} \right)^{-b} \quad (3.2)$$

where RRPOC is the rain rate,  $F$  is the export production from both surface boxes, and  $z$  is the water depth. The bathymetry of the regions of the UM and D boxes is derived from ETOPO2 ([http://www.ngdc.noaa.gov/mgg/gdas/gd\\_designagrid.html](http://www.ngdc.noaa.gov/mgg/gdas/gd_designagrid.html)). We apply  $b = 0.82$  in Eq. 3.2, which is the global average according to Berelson [2001] and also close to his estimate for the

	Sediment percentage (SD,%)	Average Martin curve value (AMC,%)
UM box	0.81	53.04
D box	2.25	12.51

Table 3.2: Sediment percentage and average Martin curve value used for the model-based estimation of benthic denitrification. “Sediment percentage” is the percentage of the surface areas of the UM and D boxes in contact with the sediment; “Average Martin curve value” represents the average of Martin curve fractions of export production reaching the sediment for each grid point of the topography data.

ETSP. An exponent of 0.4 for Eq. 3.2 in suboxic water is implied by Van Mooy et al. [2002]. Therefore, sensitivity experiments are performed with  $b = 0.4$ . From Eq. 3.2 and the fraction of the lower boundary of the respective box in contact with the seafloor, the RRPOC at the sediment surfaces of the UM and D boxes is calculated according to Eqs. 3.3 and 3.4:

$$\text{RRPOC}_{\text{UM}} = \text{EP}_{\text{U}} \times \text{SD}_{\text{UM}} \times \text{AMC}_{\text{UM}} \quad (3.3)$$

$$\text{RRPOC}_{\text{D}} = (\text{EP}_{\text{U}} + \text{EP}_{\text{S}}) \times \text{SD}_{\text{D}} \times \text{AMC}_{\text{D}} \quad (3.4)$$

where  $\text{AMC}_{\text{UM}}$  and  $\text{AMC}_{\text{D}}$  are the average Martin-curve values corresponding to the actual water depth ( $z$ ) in the grid of ETOPO2 respectively;  $\text{SD}_{\text{UM}}$  and  $\text{SD}_{\text{D}}$  represent the percentages in contact with the sediment in the UM and D boxes (Table 3.2).

### Data-based estimation of benthic denitrification

For a second and independent estimate of  $L_{\text{DIN}}$ , we combine observations from different datasets.  $\text{O}_2$  and  $\text{NO}_3^-$  concentrations for our model domain are obtained from the annual objectively analyzed mean concentrations of the WOA 2009  $1^\circ \times 1^\circ$  data [Garcia et al., 2010a,b], and interpolated over the region of our model domain to match the resolutions of the other datasets.

RRPOC is estimated from primary production following Bohlen et al. [2012]. According to the carbon-based approach of Behrenfeld et al. [2005], average annual primary production is derived from photosynthetically available radiation (PAR), the diffuse attenuation coefficient at 490 nm (K490), chlorophyll a (Chl a) and mixed layer depth (MLD). PAR, K490 and Chl a are from the Aqua-Modis satellite data (2005–2010) (<http://oceancolor.gsfc.nasa.gov/>), and MLD is from the Fleet Numerical Meteorology and Oceanography Center (FNMOC, [http://www.public.navy.mil/fltfor/cnmoc/Pages/fnmoc\\_home.aspx](http://www.public.navy.mil/fltfor/cnmoc/Pages/fnmoc_home.aspx)). Export production is estimated from primary production and sea-surface-temperature (SST) [Dunne et al., 2005],

where SST is from the WOA 2009 annual average  $1^\circ \times 1^\circ$  temperature data [Locarnini et al., 2010]. The rate of particle transport at each grid cell to the seafloor is calculated using the Martin curve (Eq. 3.2) [Martin et al., 1987]. To obtain more accurate estimates for RRPOC of our regional box model, all data processed in this experiment are interpolated on a grid of  $2' \times 2'$  in the UM box and  $20' \times 20'$  in the D box, and the ETOPO2 data ( $2' \times 2'$ ) are averaged within each  $20' \times 20'$  grid cell in the D box. The Aqua-Modis data ( $5' \times 5'$ ) and  $\text{NO}_3^-$  and  $\text{O}_2$  concentrations from WOA 2009 dataset are interpolated or averaged horizontally to match these resolutions. The vertical resolution of the  $\text{NO}_3^-$  and  $\text{O}_2$  concentrations are interpolated to resolve the bathymetry of the ETOPO2 data, and the  $\text{NO}_3^-$  and  $\text{O}_2$  concentrations closest to the sediment are applied in Eq. 3.1 for the bottom water  $\text{NO}_3^-$  and  $\text{O}_2$  concentrations.

Finally, the  $L_{\text{DIN}}$  derived from observational datasets are averaged over the regions represented by UM and D boxes to produce an annual  $\text{NO}_3^-$  loss term.

### 3.2.5 Phosphate regeneration

Phosphate regeneration is estimated according to Wallmann [2010] and Flögel et al. [2011] with both model- and data-based estimates for the rain rate. We estimate benthic  $\text{PO}_4^{3-}$  regeneration (resupply of benthic  $\text{PO}_4^{3-}$  into the water column,  $\text{Ben}_{\text{DP}}$ ) from the RRPOC degradation ratio ( $r_{\text{REG}}$ ) and the POC burial rate in the sediments (BURPOC) according to:

$$\text{Ben}_{\text{DPUM}} = \frac{\text{RRPOC}_{\text{UM}} - \text{BURPOC}_{\text{UM}}}{r_{\text{REG}}} \quad (3.5)$$

$$\text{Ben}_{\text{DPD}} = \min\left(\frac{\text{RRPOC}_{\text{D}} - \text{BURPOC}_{\text{D}}}{r_{\text{REG}}}, \frac{\text{RRPOC}}{106}\right) \quad (3.6)$$

where RRPOC is estimated with the methods described in sections 3.2.4 and 3.2.4. A minimum condition is introduced in the D-box to prevent  $\text{Ben}_{\text{DP}}$  exceeding the rain rate of particulate organic phosphate ( $\text{RRPOP} = \text{RRPOC}/106$ ) to the deep ocean, but not for the UM box, because there are possible extra sources of RRPOP, such as inputs via weathering or eolian deposition, onto the continental shelf, which is contained in the UM box in our model.

BURPOC is estimated from Eq. 3.7 for the continental shelf (UM box) and Eq. 3.8 for the deep-sea sediment (D box), and  $r_{\text{REG}}$  is the C:P regeneration ratio estimated via Eq. 3.9 following the empirical relations of Wallmann [2010].

$$\text{BURPOC}_{\text{UM}} = 0.14 \cdot \text{RRPOC}_{\text{UM}}^{1.11} \quad (3.7)$$

$$\text{BURPOC}_{\text{D}} = 0.014 \cdot \text{RRPOC}_{\text{D}}^{1.05} \quad (3.8)$$

$$r_{\text{REG}} = 123 + (-112) \times \exp\left(-\frac{\text{O}_2}{32}\right) \quad (3.9)$$

where  $\text{O}_2$  is the oxygen concentration in the ambient bottom water (in  $\mu\text{mol kg}^{-1}$ ).  $r_{\text{REG}}$  in Eq. 3.9 is higher than the Redfield ratio in oxic water, resulting in preferential P burial under oxic conditions; it is much smaller than the Redfield ratio when  $\text{O}_2 < 20 \mu\text{mol kg}^{-1}$ , indicating excess phosphate release from the sediment under suboxic conditions.

### 3.2.6 Synthesis configurations

Nitrogen deposition, benthic denitrification and phosphate regeneration are integrated into the synthesis model configurations to explore the model sensitivity to each process and their mutual interactions in the ETSP. The configurations (Syn1 to Syn4) with different benthic processes and atmospheric inputs are summarized in Table 3.1.

### 3.2.7 Model sensitivity experiments

Since our model domain only includes the top 2000 meters of the water column, the sediments only account for a small portion of the whole sediment of the ETSP (Table. 3.2). A sensitivity experiment is performed with the assumption that all of the bottom of the D box is in contact with the sediment below 500m (high benthic denitrification (high-BD), or high phosphate regeneration (high-PR)) including all  $\text{NO}_3^-$  losses by benthic denitrification and phosphate release by phosphate regeneration of the sediment.

The original work of Martin et al. [1987] and Van Mooy et al. [2002] indicate a lower value for the exponent  $b$  of Eq. 3.2 in suboxic water. We perform an additional sensitivity experiment with  $b=0.4$  according to the suggestion by Van Mooy et al. [2002], to explore the influence of benthic denitrification and phosphate regeneration under conditions of slower POC remineralization.

## 3.3 Results

### 3.3.1 Nitrogen deposition

Due to the low  $\text{NO}_3^-$  concentrations in the surface U and S boxes, the annual nitrogen input by atmospheric nitrogen deposition accounts for 63% and 10%, respectively, of nitrogen inventories of the U and S boxes. Fig. 3.2 indicates that the extra fixed-N input by nitrogen deposition reduces the growth of nitrogen fixers in the surface ocean by about  $0.72\text{--}0.73 \text{ Tg N yr}^{-1}$  (about 12% of total nitrogen fixation), which accounts for about 48% of the total fixed-N inputs into surface waters from atmospheric deposition ( $1.5 \text{ Tg N yr}^{-1}$ ). Water-column denitrification stays almost unchanged because the increase in export production (EP) of Phy is almost exactly compensated by the decrease in EP of NF, resulting in essentially unchanged total EP. As a result of the  $\sim 50\%$  of the nitrogen deposition not immediately compensated by a decline in nitrogen fixation, the model domain becomes a larger fixed-N source (Fig. 3.2). The N-loss through the lateral boundary increases from  $0.93 \text{ Tg N yr}^{-1}$  in the control configuration to  $1.7 \text{ Tg N yr}^{-1}$  in the configurations including nitrogen deposition, leading to about  $0.78 \text{ Tg N yr}^{-1}$  extra fixed-N loss from the model domain, i.e. about 50% of the total fixed-N input by nitrogen deposition. Thus, almost all the extra nitrogen input into the model domain via nitrogen deposition is offset by lower nitrogen fixation and nitrogen loss by lateral transport out of the model domain.

There is no significant influence of nitrogen deposition on the biogeochemical tracer concentrations of the model at steady state: ordinary phytoplankton (Phy) concentrations increase by 3% in the U box and even smaller changes occur in the S box, which can be attributed to the stronger nitrogen deficit in the region above the OMZ (U box) than in the open ocean (S box) (not shown). Largest changes are found for the concentration of nitrogen fixers (NF) that decrease by about 9% in the U box, counteracting the nitrogen input via nitrogen deposition (Fig. 3.3). Again in the S box, NF concentration stays almost unaltered (Fig. 3.3). There are also slight variations of the  $\text{NO}_3^-$  concentration in the UM box and of the  $\text{O}_2$  concentrations in the I and D boxes (not shown).

### 3.3.2 Benthic denitrification

The data-derived benthic denitrification and phosphate regeneration in the UM and D boxes are shown in Table 3.3. The modeled net primary production (NPP) in the surface ocean above the UM and D boxes is, respectively,  $1.4$  and  $0.87 \text{ g C m}^{-2} \text{ day}^{-1}$ , indicating higher NPP in the coastal upwelling region and lower NPP in the open ocean adjacent to the upwelling region, which is consistent with the estimate by Behrenfeld et al. [2005]. Due to the small sediment area percentages, the annual nitrogen loss by benthic denitrification is  $0.17$  and  $0.82 \text{ Tg N yr}^{-1}$  in the UM and D boxes, accounting for only about 0.14% and  $5.1 \times 10^{-3}\%$  per year, respectively,

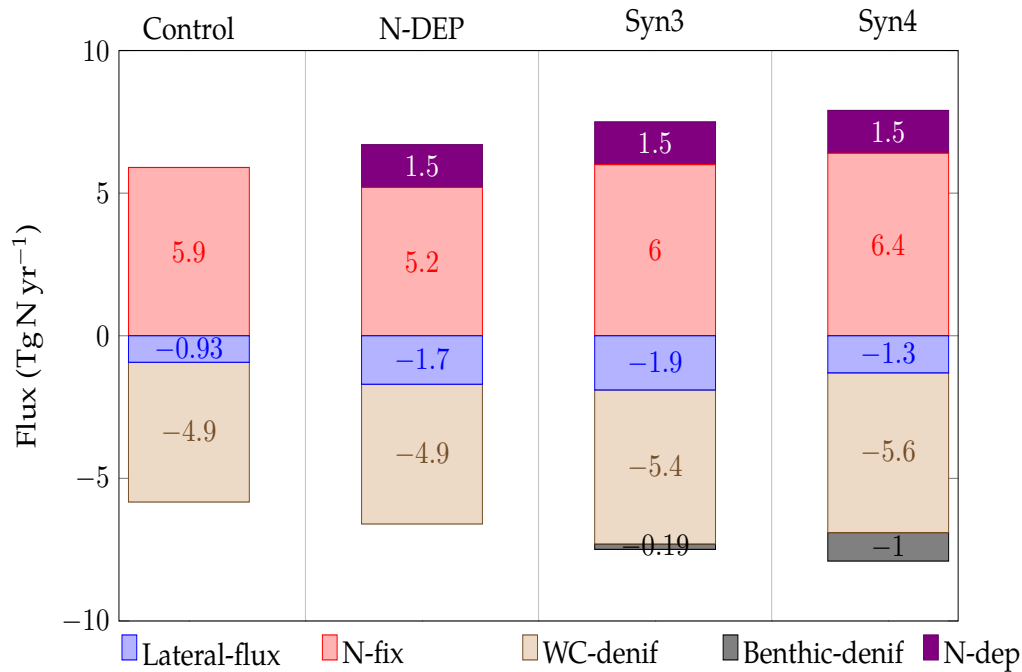


Figure 3.2: Nitrogen fluxes after including atmospheric nitrogen deposition in the control, Syn3 and Syn4 configurations defined in Table 3.1. Lateral-flux identifies the nitrogen efflux or influx through the southern boundary; N-fix represents the nitrogen fixation rate by NF; WC-denif is water-column denitrification; N-dep is the nitrogen input into surface ocean via nitrogen deposition.

	NPP	Rain rate	BD	PR
	(mg C m <sup>-2</sup> day <sup>-1</sup> )	(Tg N yr <sup>-1</sup> )	(Tg N yr <sup>-1</sup> )	(Tg P yr <sup>-1</sup> )
UM	1374.7	148.4	0.17	0.058
D	873.9	12.2	0.82	0.056
D (high-BD/PR)	873.9	12.2	8.8	0.56

Table 3.3: Fixed-N loss via benthic denitrification (BD) and phosphate release via phosphate regeneration (PR) in the UM and D boxes. High-BD indicates that the full sediment of the D box is include to estimate  $\text{NO}_3^-$  loss via benthic denitrification and phosphate release via phosphate regeneration. These values are estimated from Aqua-Modis satellite data.

of the  $\text{NO}_3^-$  inventories (Table 3.2). The higher sedimentary  $\text{NO}_3^-$  sink in the UM box can be attributed to the anoxic conditions and larger RRPOC there.

Our simulated biogeochemical tracer concentrations at steady state are quite robust with respect to including benthic denitrification (Fig. 3.3). There are only minor deviations of the MBD and DBD configurations from the control run. Nitrogen fixation rates increases by about 2.9% and 5.8% respectively in the MBD and DBD configurations (A-bars in panels MBD and

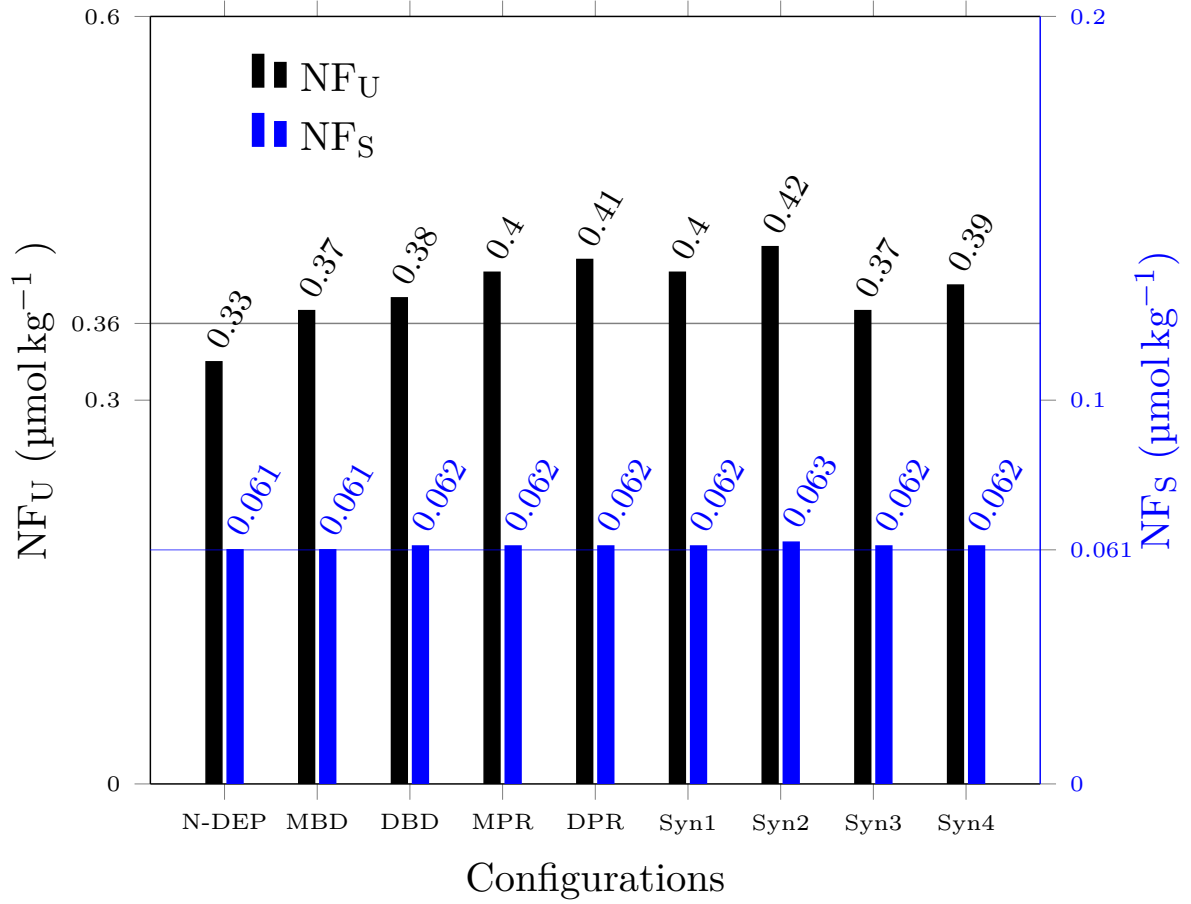


Figure 3.3: Sensitivity of simulated steady-state concentrations of nitrogen fixers  $NF_U$  and  $NF_S$  in the U and S boxes respectively. Horizontal grey and light blue lines represent the  $NF_U$  and  $NF_S$  concentrations in the control configuration respectively. Syn1, Syn2, Syn3 and Syn4 denote the “MBD+MPR”, “DBD+DPR”, “MBD+MPR+N-DEP”, and “DBD+DPR+N-DEP” synthesis configurations defined in Table. 3.1.

DBD of Fig. 3.4). Obviously, the response is stronger in the DBD configuration than in the MBD configuration, because there is an approximately 5-times larger fixed-N loss via benthic denitrification in the DBD configuration (A-bars in Fig. 3.4). The DBD configuration results in stronger responses of nitrogen fixation and lateral fluxes to benthic denitrification: the increase in nitrogen fixation can not fully compensate the nitrogen loss by benthic denitrification, thus, the model domain becomes a smaller fixed-N source, about 25% of that in the control configuration. Otherwise, the steady-state solutions of the MBD and DBD configurations are almost identical to that of the control configuration after including benthic denitrification. The temporal development of biogeochemical tracer concentrations is also insensitive to including benthic denitrification (not shown).

### 3.3.3 Phosphate regeneration

Phosphate release by phosphate regeneration accounts for about 0.23% and  $2.2 \times 10^{-3}\%$  per year, respectively, of the total phosphate inventories in the UM and D boxes (Table 3.2). The higher sedimentary  $\text{PO}_4^{3-}$  source in the UM box can be attributed to the anoxic conditions and larger RRPOC there. The phosphate release associated with benthic phosphate regeneration can stimulate nitrogen fixation and EP from the surface ocean, followed by higher water-column denitrification, owing to enhanced decomposition of exported organic matter (A-bars in MPR and DPR panels of Fig. 3.4). In the MPR configuration, nitrogen fixation and water-column denitrification increase by 12% and 11%, respectively, while in the DPR configuration, nitrogen fixation and water-column denitrification increase by about 17% and 15% (A-bars in MPR and DPR panels of Fig. 3.4). Compared with the MBD and DBD configurations, phosphate regeneration cannot turn our model domain into a smaller fixed-N source even though there is higher water-column denitrification, because nitrogen fixation can compensate for the nitrogen loss by water-column denitrification (A-bars in Fig. 3.4).

While changes in nitrogen deposition and benthic denitrification are to a large extent compensated by adjustments via simulated nitrogen fixation, phosphate is the ultimate limiting nutrient in our model domain [Su et al., 2015]. Hence the extra phosphate input into the model domain by phosphate regeneration has a more significant influence on the steady-state model results than the perturbations of the nitrogen input or loss applied in the model (Fig. 3.3). Phy concentration in the DPR configuration decreases in the U box but remains unchanged in the S box (not shown). Phy concentrations in the U and S boxes remain almost unaltered in the MPR configuration. Compared with the control configuration, NF concentrations in the U and S boxes increase by 11% and 1.6% respectively in the MPR configuration, and 14% and 1.6% respectively in the DPR configuration (Fig. 3.3). The nitrate concentration in the UM box decreases by about 4.2% in the MPR configuration and 5.2% in the DPR configuration (not shown). The temporal development of biogeochemical tracer concentrations is also robust to benthic phosphate regeneration (not shown).

### 3.3.4 Synthesis configurations

In the synthesis configurations (Tables 3.1), phytoplankton, nutrient and oxygen concentrations are quite robust with respect to the various fluxes associated with nitrogen input or removal and phosphate release from the sediment into the water column (Fig. 3.3). However, the interactions among nitrogen fixation, water-column denitrification, benthic denitrification, and phosphate regeneration result in different sensitivities of nitrogen fixation and the lateral flux to atmospheric N deposition in the presence of benthic denitrification and phosphate regeneration (Fig. 3.3). In contrast to the N-DEP configuration, nitrogen fixation rates in the Syn3 and Syn4 configurations increase by about 1.7% and 8.5% in spite of additional nitrogen in-



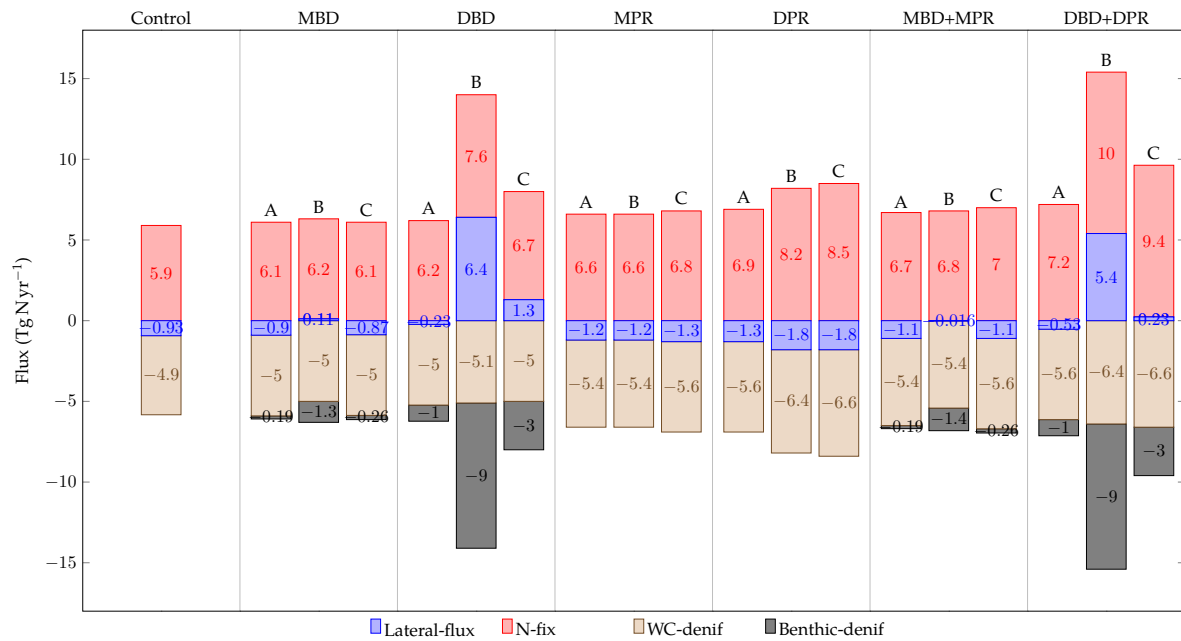


Figure 3.4: Nitrogen fluxes after including benthic denitrification or/and phosphate regeneration. Lateral-flux identifies the nitrogen efflux or influx through the southern boundary; N-fix represents the nitrogen fixation rate by NF; WC-denif is water-column denitrification; Benthic-denif represents the fixed-N loss via benthic denitrification in the model domain. Bar labels: A, main experiments; B, sensitivity experiments with high-BD; C, sensitivity experiments with Martin Curve exponent  $b=0.4$ .

put into model domain by atmospheric nitrogen deposition. The lateral flux out of the model domain ( $\text{NO}_3^-$  source) increases by about  $0.97 \text{ Tg N yr}^{-1}$  in the Syn3 configuration, which accounts for about 65% of the total nitrogen deposition, so that more than half of the extra nitrogen supplied by nitrogen deposition is not utilised locally. However, in the Syn4 configuration, the lateral  $\text{NO}_3^-$  efflux increase only accounts for about 25% of the total nitrogen deposition, with 75% of the deposited nitrogen utilised in the model domain. Less fixed-N is lost laterally from the model domain in the configurations including data-based estimates than those including model-based estimates, due to more  $\text{NO}_3^-$  loss within the model domain (Fig. 3.4). Thus, the lateral fluxes and the sensitivity to nitrogen deposition are also controlled by benthic denitrification and phosphate regeneration rather than nitrogen deposition only.

### 3.3.5 Model sensitivity

Fig. 3.5 shows the results of the sensitivity experiments with high-BD and high-PR. Compared with Fig. 3.3, the influence on the biogeochemical tracer concentrations at steady state is stronger, due to the larger  $\text{NO}_3^-$  loss via benthic denitrification and  $\text{PO}_4^{3-}$  release via phosphate regeneration (Table. 3.3). High-BD or high-BD together with high-PR can even turn our model

domain into a  $\text{NO}_3^-$  sink (B-bars in panels DBD and DBD+DPR of Fig. 3.4).

Applying the Martin Curve exponent  $b=0.4$  also amplifies the influence of benthic denitrification and phosphate regeneration on phytoplankton and biogeochemical tracers, although the effect is weaker than in the high-BD and high-PR configurations. For example,  $\text{NF}_U$  increases by as much as 33% in the DBD+DPR configuration, and  $\text{NF}_S$  increase about 34% in the DBD configuration (Fig. 3.6). Compared with A-bars in Fig. 3.4, this enhanced influence results from the higher  $\text{NO}_3^-$  loss through benthic denitrification and phosphate input via phosphate regeneration (C-bars in Fig. 3.4).

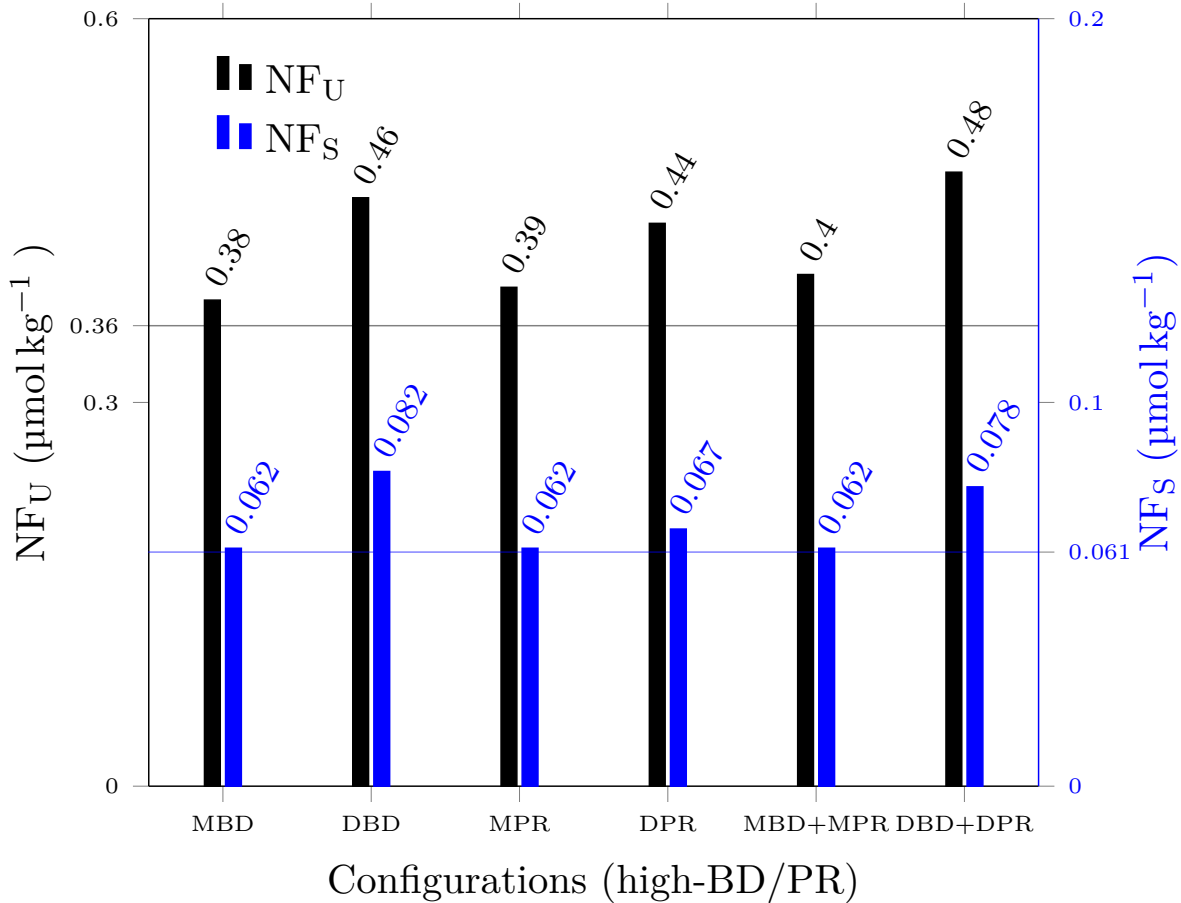


Figure 3.5: Sensitivity of simulated steady-state concentrations of nitrogen fixers ( $\text{NF}_U$  and  $\text{NF}_S$ ) in the U and S boxes respectively after incorporating high-BD and high-PR. Horizontal grey and light blue lines represent the  $\text{NF}_U$  and  $\text{NF}_S$  concentrations in the control configuration.

Due to the higher RRPOC reaching the sea floor below the water column with suboxic conditions, benthic denitrification increases by about 42% and 198% (A- and C-bars of panels MBD and DBD in Fig. 3.4) and phosphate regeneration increases about 36% and 200% respec-

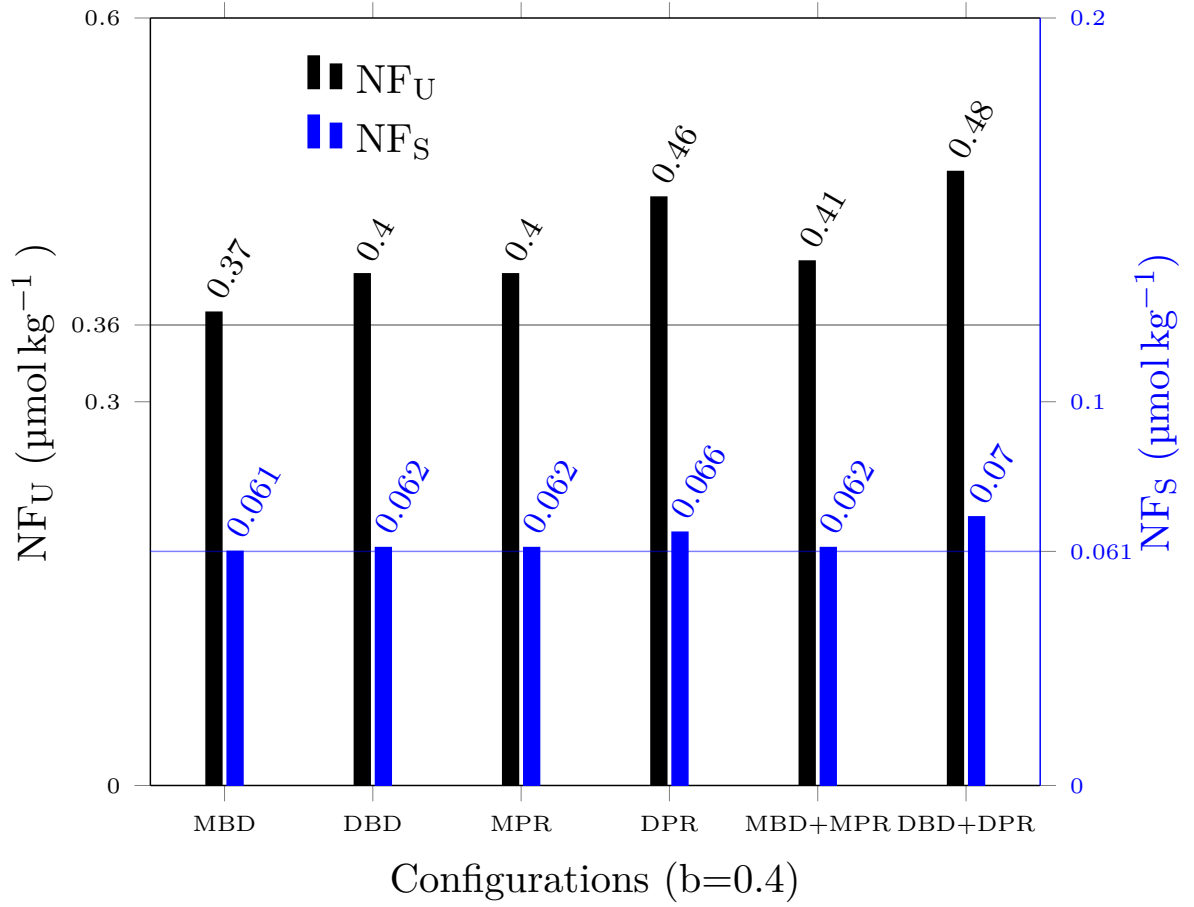


Figure 3.6: Sensitivity of simulated steady-state concentrations of nitrogen fixers ( $NF_U$  and  $NF_S$ ) in the U and S boxes respectively after applying  $b=0.4$  for Eq. 3.2. Horizontal grey and light blue lines represent the  $NF_U$  and  $NF_S$  concentrations in the control configuration.

tively in model- and data-based estimations in the sensitivity experiments with Martin Curve value  $b=0.4$ . Our model domain switches to a  $NO_3^-$  sink in the DBD and DBD+DPR configurations with  $b=0.4$  (C-bars in Fig. 3.4). Comparing A- and C-bars of panel DBD in Fig. 3.4, we find that higher benthic denitrification can stimulate nitrogen fixation, but the water-column denitrification remains constant. However, comparing A- and C-bars of panel DBD+DPR in Fig. 3.4, we find that higher benthic denitrification can increase nitrogen fixation and water-column denitrification, indicating important role of  $PO_4^{3-}$  in the balance nitrogen inventory. This shows a positive feedback between water-column denitrification in the OMZ and benthic denitrification below, caused by slower remineralisation under anoxic conditions, which results in more RRPOC reaching the sea floor. All above comparisons indicate that phosphate limitation could be responsible for breaking this positive feedback under the assumption of our model that  $PO_4^{3-}$  is the only limiting factor for the growth of nitrogen fixers.

### 3.4 Discussion and conclusions

The impact of nitrogen deposition on the ETSP has rarely been investigated so far, since this region is believed to receive less atmospheric fixed-N deposition compared with the coasts of western Europe, South and East Asia [Dentener et al., 2006, Duce et al., 2008]. The influence of anthropogenic nitrogen deposition on the biogeochemical cycles of the open ocean is increasing and the increase in atmospheric nitrogen deposition will probably induce an approximately 10% rise in carbon sequestration on land and in the ocean by 2030 [Duce et al., 2008, Reay et al., 2008]. The ETSP, a typical N-deficit region due to denitrification in the OMZ, is likely to be sensitive to anthropogenic nitrogen deposition. We find that, in our model, nitrogen deposition can inhibit  $N_2$  fixation by relieving nitrogen limitation for Phy: this counteracts the effect of atmospheric nitrogen input. This is in line with the finding that  $N_2$  fixation decreases with increasing nitrogen deposition in global-scale models that use essentially the same assumptions about the environmental controls on marine nitrogen fixation. [Krishnamurthy et al., 2007, 2009, 2010, Zamora et al., 2010]. Another portion of the deposited nitrogen is exported out of the model domain since not all the deposited nitrogen can be taken up by Phy locally because of phosphate limitation (Fig. 3.2). Appendix A shows the model results when there is facultative  $N_2$  fixation, which slightly enhances the strength of the negative feedback between nitrogen fixation and nitrogen deposition.

In a 3-D biogeochemical model study of the nitrogen fixation response to benthic denitrification, Somes et al. [2013] found that an increase of benthic denitrification can stimulate  $N_2$  fixation, but this occurred under the condition that  $N_2$  fixation was tuned to compensate for the fixed-N loss by reducing mortality and predation rates. Nitrogen fixation can also be stimulated by a N-deficit [Deutsch et al., 2007], which can result from benthic denitrification. In our 2-D model, we also find that nitrogen fixation can be enhanced by the N-deficit caused by benthic denitrification, so that nitrogen fixation can somehow compensate for the nitrogen loss.

We find a strong increase in primary production after incorporating benthic phosphate regeneration, which is mainly attributed to nitrogen fixation (Panels MPR and DPR in Fig. 3.4). Phosphate regeneration will be enhanced under  $O_2$  deficit conditions, and the enhanced phosphate release will stimulate primary production, finally resulting in the expansion of OMZs and a possible positive feedback loop leading to more benthic phosphate regeneration [Van Cappellen and Ingall, 1994, Wallmann, 2010]. However, our model domain only represents the upper 2000 meters of the ocean and its sediments only account for a small fraction of total sediment in the ETSP. The model results incorporating benthic denitrification and phosphate regeneration for a case where all of the bottom of the D box is assumed in contact with the sediment are shown in Figs. 3.4 and 3.5. For our parameterization of nitrogen fixation being favoured in N-deficit waters, the increase of water-column denitrification can always be compensated by increased nitrogen fixation when there is only phosphate regeneration (panels

MPR and DPR in Fig. 3.4). Our model domain can only turn into a fixed-N sink after including high-BD of the full sediment.

Due to the simplicity and computational efficiency of our box model, it is relatively easy to explore model sensitivity to processes related to the nitrogen budget of the ETSP. Even though some spatial and temporal variations are missing compared with results from the global circulation models [Krishnamurthy et al., 2007, 2010, Zamora et al., 2010], we can efficiently diagnose the regional impacts at steady state. Stimulatory effects can be identified, respectively, between nitrogen fixation and water-column denitrification, phosphate regeneration and nitrogen fixation, phosphate regeneration and water-column denitrification, and atmospheric deposition and lateral  $\text{NO}_3^-$  transport (Fig. 3.4). Depressive effects occur between atmospheric deposition and nitrogen fixation, benthic denitrification and the lateral  $\text{NO}_3^-$  transport (Fig. 3.4). The model sensitivity to processes related to the nitrogen budget of the OMZ in the ETSP is illustrated in Fig. 3.7. The Nitrogen fixation rate can be enhanced by benthic denitrification, compensating for part of the  $\text{NO}_3^-$  loss. The stimulatory effect between nitrogen fixation and water-column denitrification can help to keep a fixed-N balance in the ocean. The extra fixed-N input by nitrogen deposition can be counteracted by decreased nitrogen fixation and partially removed by lateral flux. All of these local responses combined constitute a nitrogen-balancing mechanism in the ETSP. In the high-BD sensitivity experiment, our model domain turns into a  $\text{NO}_3^-$  sink (Fig. 3.4). The  $\text{NO}_3^-$  inventory in the ETSP is determined by nitrogen fixation, water-column denitrification, benthic denitrification and lateral  $\text{NO}_3^-$  flux. Since nitrogen fixation and water-column denitrification ensue only in the surface ocean and the OMZ, which are included in our model domain, we cannot rule out that the ETSP including sedimentary denitrification is a  $\text{NO}_3^-$  sink, which is consistent with many model-derived or observational results [Ganachaud and Wunsch, 2002, Kalvelage et al., 2013]. Extra phosphate input into the model domain via phosphate regeneration can increase water-column denitrification significantly due to the increase of EP from the surface ocean. However, phosphate regeneration alone can not turn our model domain into a  $\text{NO}_3^-$  sink, which corroborates that the water column of ETSP is net source of  $\text{NO}_3^-$ .

The remineralization rate of organic matter is thought to be reduced under anoxic conditions [Martin et al., 1987, Van Mooy et al., 2002], which results in more RRPOC reaching the sediments. According to the analysis of Bohlen et al. [2012], benthic denitrification is very sensitive to RRPOC, i.e., higher RRPOC results in higher benthic denitrification. Based on the findings that higher benthic denitrification can increase nitrogen fixation, higher nitrogen fixation could result in higher water-column denitrification and the expansion of OMZs and finally a positive feedback between water-column and benthic denitrification. But this positive feedback is only observed in the configurations with phosphate input via phosphate regeneration, which indicates that  $\text{PO}_4^{3-}$  limitation could prevent this positive nitrogen loss feedback.

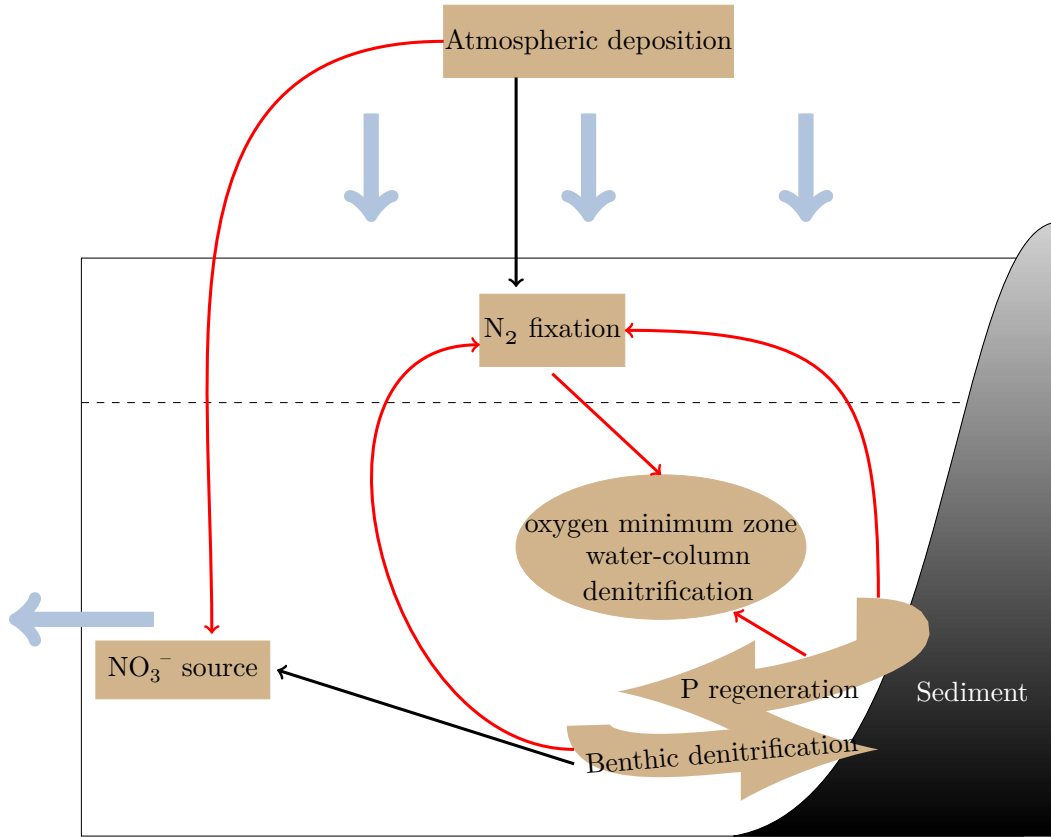


Figure 3.7: Schematic of the model sensitivity to different processes related to the nitrogen budget of the ETSP. The red solid lines present stimulatory effects, and the black solid lines represent depressive effects.

### 3.5 Appendix A

It is known from laboratory experiments that diazotrophic phytoplankton can also utilize DIN for growth [e.g., Holl and Montoya, 2005]. In contrast to our NF model where NF can only fix N<sub>2</sub> from the atmosphere, Schmittner et al. [2008] introduced a formulation to allow also NO<sub>3</sub><sup>-</sup> uptake by diazotrophs. In Schmittner et al. [2008]’s model, nitrogen fixers preferentially use nitrate when available and cover only the residual N demand by N<sub>2</sub> fixation, denoted as facultative N<sub>2</sub>-fixation. Thus, we explore the effect of facultative N<sub>2</sub>-fixation on our model results with extra fixed-N input by nitrogen deposition.

Compared to results from the configurations in which NF can only fix N<sub>2</sub> (Fig. 3.3), both Phy and NF in the U-box are more robust to the extra nitrogen input via nitrogen deposition, for instance, Phy decreases by 1.5% (facultative N<sub>2</sub>-fixation) vs 2.9% (N<sub>2</sub> fixation), and NF increases by 2.5% (facultative N<sub>2</sub>-fixation) vs 11% (N<sub>2</sub> fixation). Again, the biogeochemical concentrations at steady state are relatively insensitive to nitrogen deposition (not shown).

There is a stronger negative feedback between nitrogen deposition and facultative N<sub>2</sub>-fixation,

since nitrogen fixation is reduced by about 21% (facultative  $\text{N}_2$ -fixation) compared to 12% (obligate  $\text{N}_2$  fixation) (Fig. 3.8). The lateral fluxes of  $\text{NO}_3^-$  only accounts for about 34% of the extra nitrogen input by nitrogen deposition (facultative  $\text{N}_2$ -fixation) compared to 50% (obligate  $\text{N}_2$  fixation) (Fig. 3.8). Thus, facultative  $\text{N}_2$ -fixation can adjust nitrogen fixation in response to nutrient concentrations in the surface boxes and control the magnitude of our model domain being a  $\text{NO}_3^-$  source.

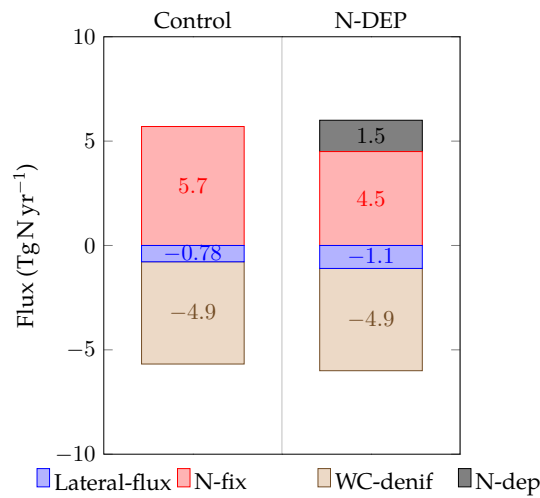


Figure 3.8: Nitrogen fluxes after including atmospheric nitrogen deposition in the model with facultative  $\text{N}_2$ -fixation. Labels are the same as those in Fig. 3.2.

### 3.6 Acknowledgements

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## Chapter 4

# **Coupled physical/biogeochemical modeling of the oxygen minimum zone in the eastern tropical South Pacific: sensitivity to the southern boundary conditions**

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This is a manuscript in preparation.

**Abstract** The impact of the southern boundary on the Oxygen Minimum Zone (OMZ) of the eastern tropical South Pacific (ETSP) is investigated with a high-resolution coupled physical-biogeochemical model. Model assessment of the realistic model configuration suggests good agreement of horizontal and vertical oxygen and nutrient distributions with observations. Comparisons among different model configurations indicate that the southern boundary is an important oxygen source to the OMZ in the ETSP, mainly influencing the southern border and the coast of the region. Lateral flux estimations show that our model domain is gaining oxygen, but losing nitrate and phosphate through the southern boundary. The southern boundary is found to have a more significant influence on  $O_2$  concentration of the deep layer than that of the upper layer, suggesting a pathway of deep northward intrusion of well-oxygenated southern waters. The Southern Ocean is thought to be sensitive to global warming due to reductions in deep convection, resulting in a significant decrease in  $O_2$  concentrations. This study can give some insights into the impacts of Southern Ocean variations on the  $O_2$  cycle of the ETSP.

## 4.1 Introduction

The Humboldt Current System, known as the most productive eastern boundary current system, is dominant at the surface and subsurface of the eastern South Pacific. In the northern part of the Humboldt Current System, the Peru Current System (PCS) is located at the eastern tropical South Pacific (ETSP) off Peru, which hosts one of the most significant upwelling regions and permanent oxygen minimum zones (OMZs) of world ocean [Hellya and Levin, 2004, Paulmier and Ruiz-Pino, 2009]. In the OMZ,  $NO_3^-$  is converted to  $N_2$  or  $N_2O$  by canonical denitrification or anammox, resulting in a nitrate deficit relative to phosphate [Codispoti and Christensen, 1985]. Thus, the ETSP is a location for active reactions associated with nitrogen cycle, and has significant impacts on local and even global nitrogen cycles.

The ETSP has a rich water mass structure, with water sources mainly from the equatorial Pacific and the Southern Ocean to feed waters of different depths. The surface of the PCS is the equatorward Peru Coastal Current (PCC), with a width of 500-1000 km from the coast [Strub et al., 1998]. PCC is directly associated with the coastal upwelling, thus featured by cold and salty water [Stramma et al., 2010]. It carries the subtropical water (STW) and subantarctic water (SAAW) with low  $NO_3^-$  and  $PO_4^{3-}$  concentrations, and feeds the South Equatorial Current (SEC), which is the westward surface current of the Equatorial Current System (ECS) [Llanillo et al., 2013, references therein]. The Peru-Chile Under Current (PCUC) is a poleward flow below the PCC, which dominates the subsurface layer and extends from 50m to 700m along the western coast of South America. Its depth increases toward south and transports salty equatorial subsurface water (ESSW), which is rich in nutrients and extremely low in  $O_2$  [Wyrki, 1963, Penven et al., 2005, and references therein]. Since the PCUC is thought to be one of the

major sources of the upwelled water of the ETSP, the oxygen and nutrient concentrations of it are important for the study of the upwelling region and the OMZ below [Penven et al., 2005, and references therein]. As another surface current belonging to the PCS, the offshore Peru Oceanic Current (POC) extends from the surface to a depth of 700m and veers westward around 15–20°S [Wyrtki, 1963, Penven et al., 2005]. Offshore of PCC and PCUC, the Peru-Chile Counter Current (PCCC) is featured by warm water originating from the split of Equatorial Under Current (EUC), and ranges from 80°W to 84°W and 8°S to 35°S [Huyer et al., 1991, Penven et al., 2005]. The PCCC flows southward carrying ESSW and reaches a maximum speed of  $0.1\text{ m s}^{-1}$  at 50m depth [Huyer et al., 1991].

Due to its remarkable characteristic of the PCS being in proximity and directly connected to the equatorial ocean at its northern border both in the surface ocean (0–50m) and below the surface, the ECS is one of the water sources of the PCS. PCUC is thought to originate from the eastward Equatorial Under Current (EUC), which splits when arriving at the American landmass. Both the primary South Subsurface Countercurrent (sSSCC) and secondary South Subsurface Countercurrent (pSSCC) could be complementary sources of the PCUC [Lukas, 1986, Montes et al., 2010]. The connection between the PCUC and the equatorial eastward subsurface currents (EUC, pSSCC, sSSCC) is very important for understanding the formation of the OMZ in the ETSP. However, Montes et al. [2010] find in a high-resolution regional model that only about 30% of the PCUC is fed by the three subsurface equatorial currents (EUC, pSSCC, sSSCC), among which only the two SSCCs contribute substantially. The remaining part of the PCUC comes from an alongshore recirculation associated with flows below and from the southern part of the region (south of 9°S) [Montes et al., 2010], indicating the southern border being important for the formation and maintenance of the OMZ. The box model of Su et al. [2015] for the ETSP also shows that the  $\text{O}_2$  ventilation from the southern boundary is very important in maintaining a realistic  $\text{O}_2$  concentration in the deep ocean and the water column of the ETSP is a source of  $\text{NO}_3^-$  and  $\text{PO}_4^{3-}$  under its parameterization.

There are mainly two water masses in the intermediate and deep levels of the ETSP: antarctic intermediate water (AAIW) and pacific deep water (PDW). The new-found Peru-Chile Deep Coast Current (PCDCC) flows below the PCUC and carries AAIW between 700m and 1200m [Chaigneau et al., 2013]. Typically low in nutrients and high in dissolved  $\text{O}_2$  concentrations, the northward AAIW brings oxygen into the ETSP [Schneider et al., 2003, and references therein]. Below AAIW sits the PDW, which is a vast water mass flowing southward. The PDW is thought to be homogenous and the oldest water mass in the ocean, which carries water with low  $\text{O}_2$  but high nutrient concentrations [Reid, 1997, Tomczak and Godfrey, 2003, Llanillo et al., 2013].

In this study, the impact of the southern boundary on the OMZ characteristics and the nutrient distributions of the ETSP is investigated in a high-resolution coupled physical-biogeochemical model. The model is set up in a climatological configuration similar to Echevin et al. [2008] and Montes et al. [2010], but with a smaller domain in order to exclude the impacts of ECS and fo-

cus on the southern boundary. The model assessments in section 4.2.5 show that our model can well reproduce both physical and biogeochemical features of this region, which provides the precondition for studying the southern boundary conditions. Then we investigate the impact of different southern boundary conditions on the OMZ and the whole ETSP in terms of (1) OMZ structure evolution, (2)  $O_2$  and nutrient contents, (3)  $O_2$  and nutrient distributions, and (4) lateral fluxes. We find that the southern boundary conditions have a more significant influence on the deep than on the surface ocean. The ETSP is gaining  $O_2$  from the southern boundary, but losing both  $NO_3^-$  and  $PO_4^{3-}$ .

## 4.2 Methods

To investigate the role of the southern boundary on the OMZ off Peru, a coupled physical and biogeochemical model to simulate the dynamics of this region is applied. The physical model employed is the regional model ROMS (Regional Oceanic Model System). This model is coupled with the biogeochemical model PISCES (Pelagic Interaction Scheme for Carbon and Ecosystem Studies).

### 4.2.1 ROMS oceanic model

ROMS is a free surface and terrain-following vertical coordinate model, which is designed to enhance resolution near the sea surface [Shchepetkin and McWilliams, 2005, 2009]. As described by Shchepetkin and McWilliams [2005], this model employs a mode-splitting algorithm to separate the time-scales of barotropic and baroclinic processes, by solving the vertically-integrated barotropic momentum equations [Shchepetkin and McWilliams, 2009]. The unresolved processes involved in vertical mixing are parameterized using a non-local K-Profile (KPP) introduced by Large et al. [1994]. Its open boundary conditions provide outward radiations and nudging towards prescribed external boundaries (sponge layers). Explicit lateral viscosity is null everywhere in the simulated model domain, except in the sponge layers at open boundaries.

### 4.2.2 PISCES model

PISCES is a biogeochemical model which simulates the marine biological productivity and describes the biogeochemical cycles of carbon and the main nutrients (N, P, Si, Fe) [Aumont and Bopp, 2006]. It includes two phytoplankton (nanophytoplankton and diatoms), and two zooplankton (microzooplankton and mesozooplankton) size classes. The PISCES model considers phytoplankton growth limitation by five different nutrients:  $NO_3^-$ ,  $NH_4^+$ ,  $PO_4^{3-}$ ,  $SiO_4^{3-}$  and Fe. There are three non-living compartments: semi-labile dissolved organic matter, small

and large sinking particles. The model also simulates dissolved inorganic carbon, total alkalinity and dissolved oxygen [Aumont and Bopp, 2006]. All parameters for this region follow Echevin et al. [2014].

The nitrogen-related biogeochemical processes in PISCES include nitrogen fixation, denitrification, nitrification and ammonification [Aumont, 2005]. Diazotrophic phytoplankton can fix  $N_2$  under the conditions that: 1) water temperature is above 20 °C, 2) there is insufficient fixed-N in the environment, 3) iron is available. Nitrogen fixation is restricted to the sea surface and annual total nitrogen fixation should balance denitrification to ensure nitrogen conservation in the model domain. Denitrification ensues only when oxygen concentration is less than 6  $\mu\text{mol kg}^{-1}$ . Nitrification represents the conversion of  $\text{NH}_4^+$  to  $\text{NO}_3^-$  due to bacterial activity, which is assumed to be photoinhibited and  $\text{O}_2$  concentration dependent. Ammonification is responsible for the remineralization of particulate organic matter and dissolved organic matter to  $\text{NH}_4^+$ , which can be utilised directly by phytoplankton again.

### 4.2.3 Model configurations

#### The standard configuration

To capture the role of the southern boundary on the OMZ off Peru, we implement a regional configuration centered at the core of the eastern tropical south Pacific OMZ. This configuration allows a close comparison with the previous study by Su et al. [2015], which suggests an important role of open boundaries on preventing  $\text{O}_2$  depletion in the deep ocean of the ETSP. The present configuration has a rectangular grid extending from 4°S to 16°S in latitude and from 72°W to 91°W in longitude. The horizontal grid has an isotropic resolution of  $1/9^\circ$ , corresponding to about 12 km, and contains 172x111 grid points. The vertical grid has 32 vertical sigma-levels, parameterized to allow good representation of upper layer ocean processes ( $\text{theta}_s=6$ ,  $\text{theta}_b=0$ ,  $\text{hc}=10$ ). Thus, the vertical resolution varies according to the water column depth, from a minimum of 37.4cm to a maximum 5.04m for the surface layer and from 10.43m to 776.71m for the bottom layer.

#### The idealized configurations

The role of the southern boundary on the OMZ off Peru is investigated by comparing simulations with different southern boundary settings, from completely closed boundary conditions to removing all the incoming oxygen and nutrient supplies into model domain through the southern boundary. There are four configurations in total developed in this study: (1) STD is the standard configuration introduced above. (2) CLOSEBOUND is identical to the STD simulation except that the southern boundary is totally closed, suppressing the exchange of water masses properties with the southern subtropical Pacific. (3) NOO2 is identical to the STD simulation except that there is no oxygen supply into the model domain via advection from the southern boundary. (4) NOO2NO3PO4 is identical to the NOO2 simulation except that there are no  $\text{NO}_3^-$  and  $\text{PO}_4^{3-}$  supplies into the model domain via advection from the southern

boundary. We consider the STD configuration the most realistic, and the others as the idealised experiments.

### Simulations

All simulations of the four model configurations are conducted over 30 climatological years, with model outputs averaged and stored every 3 days. The simulated physical and biogeochemical dynamics reach an equilibrium after 25 years (Appendix A). Therefore, we have constructed the mean dynamics (monthly, seasonal and annual) from the last five years of the model simulation (years 26–30).

#### 4.2.4 Data sets used for open boundary conditions, forcing and model assessments

The model grid, forcing, initial and boundary conditions are built with the ROMSTOOLS package [Penven et al., 2008]. The bottom topography and coastline of our model domain are derived from the ETOPO2 dataset [Smith and Sandwell, 1997], which provides a fine 2' resolution to avoid potential errors in the horizontal pressure gradient. COADS ocean surface 0.5° resolution monthly climatology data are applied for the model heat and fresh water fluxes at the ocean surface [Silva et al., 1994]. For surface wind stress, the QuickSCAT satellite scatterometer data, which is gridded at a resolution of 0.5° and calculated over 2000–2007, is employed [Liu et al., 1998]. Pathfinder, a finer SST climatology (1/12°), is employed to replace the coarser SST dataset [Casey and Cornillon, 1999]. The initial values and boundary conditions of the biogeochemical tracers including  $O_2$ ,  $NO_3^-$ ,  $PO_4^{3-}$ , and  $SiO_4$  are interpolated from CISRO Atlas of Regional Seas [CARS, 2009, <http://www.cmar.csiro.au/cars>] to the model resolution. The dynamical variables (temperature, salinity and velocity fields) for ROMS open boundary conditions are provided by the monthly climatology computed from the Simple Ocean Data Assimilation (SODA) reanalysis [Carton and Giese, 2008]. The interannual variabilities such as EL-Niños are explicitly excluded.

AVISO satellite altimetric data (<http://www.aviso.oceanobs.com/>) in the period from January 1990 to December 2010 are applied to evaluate the sea surface height variability. The CISRO Atlas of Regional Seas (Ridgway et al. [2002], Dunn) monthly climatological data with 1/2° resolutions for nutrients, oxygen, salinity and temperature are used for comparison with model outputs to assess our model skills. The monthly climatological SeaWiFS products of level 3-binned data for Chlorophyll(Chla) over 2005–2010 are used for comparison with simulated Chla concentration at surface, which is the sum of diatom and nanophytoplankton chlorophyll.

#### 4.2.5 Model assessment

The simulated physical and biogeochemical dynamics from the STD configuration are compared with existing observations and previous studies, in order to evaluate the realistic repro-

duction of the dynamics of this region.

Figure 4.1 shows the annual mean sea surface height from the AVISO data and the STD configuration. From this dynamic, the Peru coastal current (PCC) along the coast and the equatorward flow of the Peru Oceanic current (POC) offshore are visible. Close to the southern border, between 12 and 15°S, the westward current represents the Peru Oceanic Current (POC). Thus, our simulation can reproduce the general patterns of surface currents compared with those estimated from the AVISO data.

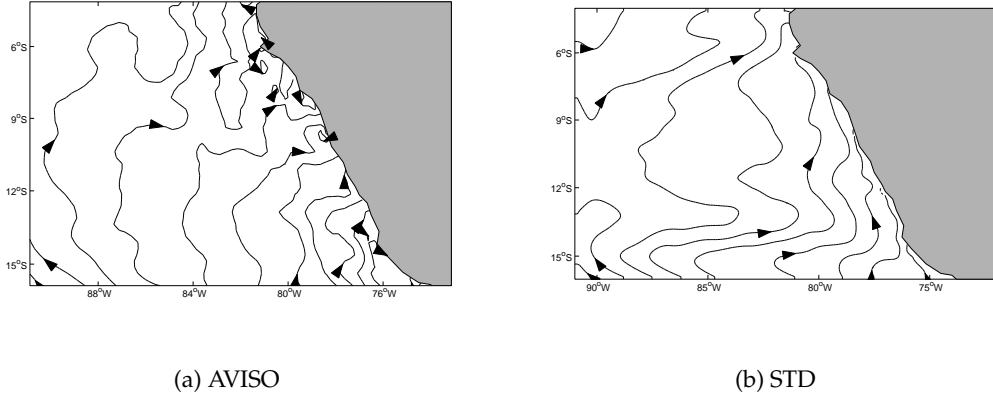
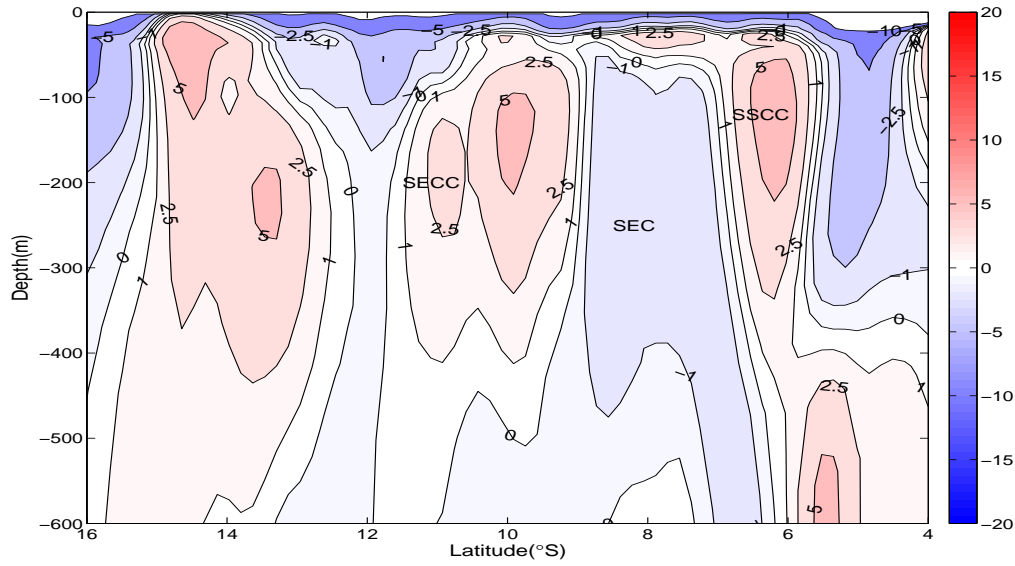
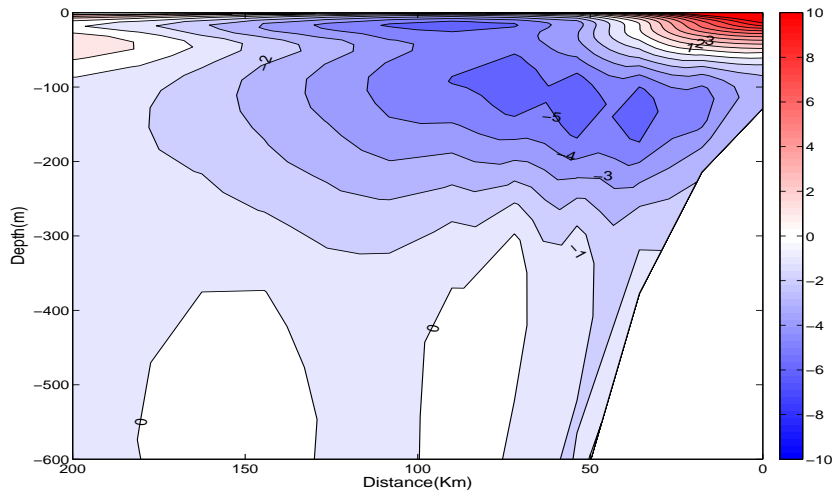


Figure 4.1: Surface currents derived from the annual mean sea surface height for both data (a) and the STD configuration (b).

The vertical structures of the zonal and meridional velocities (Fig. 4.2) also suggest good agreement with the previous observations and model results [Penven et al., 2005, Montes et al., 2010, Czeschel et al., 2011]. The simulated westward South Equatorial current (SEC) separates the eastward flow of the subsurface counter currents (SSCC) and the South Equatorial Counter-current (SECC) of this region, in agreement with the previous observations [Czeschel et al., 2011] and model studies [Montes et al., 2010]. The eastward flow of the SSCC is centered at around 6°S in the simulated results as documented in the observational study by Czeschel et al. [2011]. Further south, at around 12°S, the structure of the SECC is well represented in the model simulation [Czeschel et al., 2011]. Along the coast, the northward PCC is dominant at the upper layer of the onshore side, and overlaying the core of the Peru Chile Undercurrent (PCUC), which is in agreement with the previous studies [Penven et al., 2005, Montes et al., 2010, Echevin et al., 2014]. The subsurface PCUC flows poleward with the velocities up to  $10 \text{ cm s}^{-1}$ , in agreement with the previous studies [Penven et al., 2005, Montes et al., 2010, Echevin et al., 2014].



(a) u-86°W-Feb



(b) v-12°S

Figure 4.2: Vertical sections of zonal and meridional velocities ( $\text{cm s}^{-1}$ ). (a) mean zonal velocity at 86°W in February; (b) annual mean meridional velocity at 12°S.

A Taylor diagram is shown in Fig. 4.3 to evaluate the modeling abilities of the STD configuration in terms of both physical (temperature and salinity) and biogeochemical (oxygen, nitrate, phosphate and surface chlorophyll) fields in an annual mean time scale [Taylor, 2001]. The simulated fields of the STD configuration are compared with climatological CARS 2009 data in statistical metrics of the standard deviation, the correlation coefficient and root mean square (RMS). For the physical variables, correlation coefficients are above 0.97 for both surface and 100-600m depth of the model domain. The normalized standard deviations are very close to 1.0 and the centered RMS differences are between 0.2 and 0.3 (Fig. 4.3). The simulated bio-



geochemical variables are also in agreement with the CARS 2009 and SeaWiFS observations, with a correlation coefficient above 0.7. The RMS and standard deviation are more dispersive. The surface chlorophyll and nitrate in the ocean interior show low variations compared to observed fields. The variations in oxygen (both at surface and in ocean interior), surface nitrate and phosphate in the ocean interior are larger than observations (Fig. 4.3).

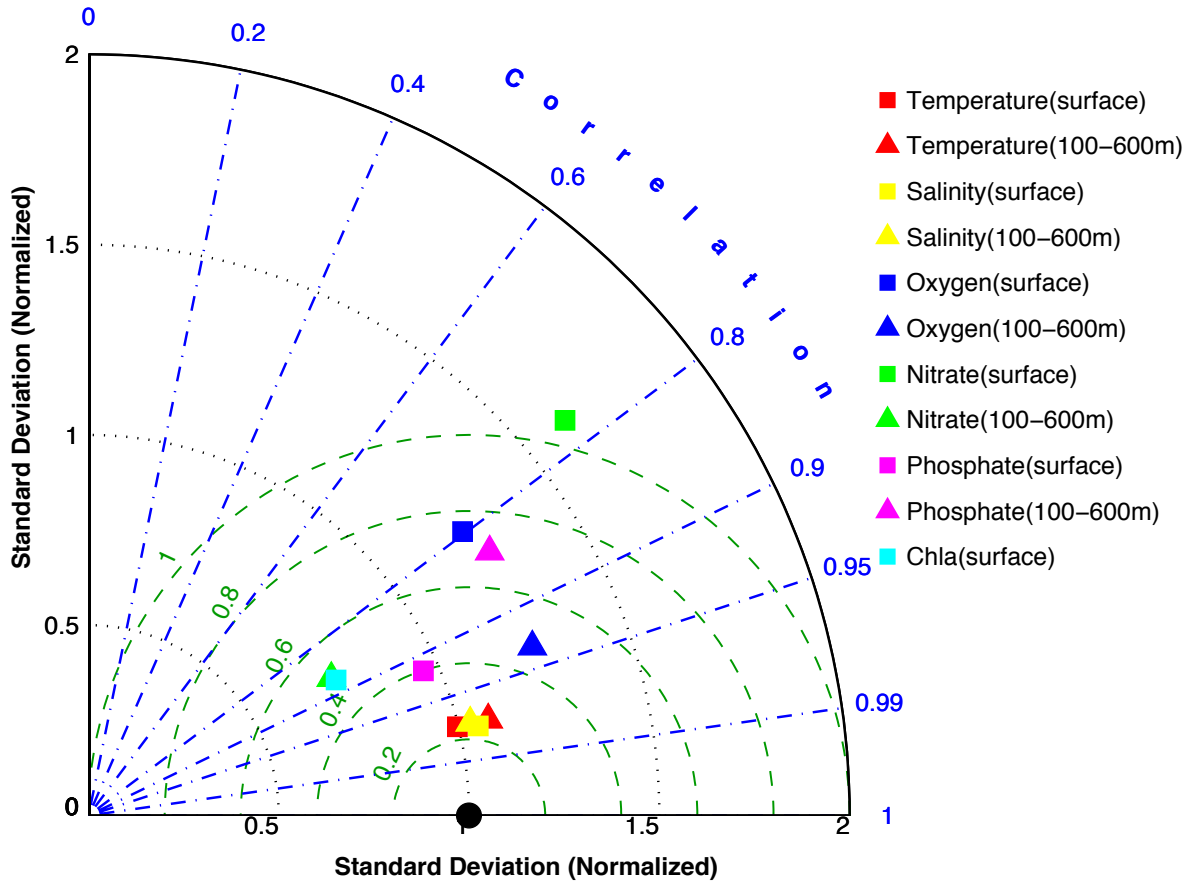


Figure 4.3: Taylor diagram for annual average temperature (red), salinity (yellow), oxygen (blue), nitrate (green), phosphate (magenta) and chlorophyll (cyan) concentrations. The radial distance from the origin (the black filled circle) is proportional to the standard deviation, which is normalized by the standard deviation of the observational data. The correlation between the model and data is shown by the angular coordinates (blue dash-dot lines). The distance of each model point from the origin represents the root-mean-square (RMS, green dashed lines). The statistics are performed for the horizontal surface slice and 100-600m depth of the model domain (to represent the core of OMZ) with both model outputs and data to evaluate the model skills. The model outputs are the last 5 years annual means; the data are from the CARS 2009 annual climatologies.

The vertical structures of simulated temperature and salinity also suggest a realistic representation of the dynamics of this region. In the upper surface layer (Figs. 4.4 and 4.5), the warmer and salty subtropical waters (STW) dominates the water mass properties in both off-

shore and coastal sections. Along the coast, the relatively low temperature and salinity suggest the upwelled subsurface waters induced by the equatorial winds along the coast ((a) and (b) in Figs. 4.4). The simulated subsurface waters are colder and fresher, characteristic of the equatorial subsurface water (ESSW). The ESSW is also characterized by extremely low oxygen and high nutrient concentrations ((c), (d) and (e) in Figs. 4.4 and 4.5). Above the less oxygenated ESSW, the simulated STW is well oxygenated and with low nutrient, which is in agreement with observation ((c) in Figs. 4.4 and 4.5). The only deficiency in the meridional section is that both temperature and salinity are slightly higher than data in the top 100m between 12 and 15°S ((a) and (b) in Fig. 4.5).

### 4.3 Results and discussion

To gain insights into the influence of southern boundary on the ETSP dynamics, we analyse the evolution of OMZ structures, the  $O_2$  and nutrient contents, the volume distributions as a function of  $O_2$  and nutrient concentrations, and the vertical profiles of average  $O_2$ ,  $NO_3^-$  and  $PO_4^{3-}$  concentrations of the model domain of the four configurations defined in section 4.2. At the end, the lateral fluxes of  $O_2$  and nutrients for our model domain are examined.

#### 4.3.1 The effect of southern boundary on the OMZ structure

The structures of the OMZ (defined as  $O_2$  concentration  $< 20 \mu\text{mol kg}^{-1}$ ) derived from the oxygen distribution of both realistic STD and idealized simulations are presented in Fig 4.6. The structure of the OMZ in the STD configure is comparable with that estimated from the CARS 2009 data, with extended OMZ thickness close to the continental shelf and reduced thickness in direction toward the southern border ((a) and (b) in Fig. 4.6). The maximum OMZ thicknesses are about 600m in both the estimate with CARS 2009 data and the STD configuration.

In the CLOSEBOUND configuration, the low-oxygen waters are vertically expanded in the southern part of the domain, with maximum thickness of 700m ((c) in Fig. 4.6). This expansion is pronounced in the south-western side of the domain, suggesting an important role of  $O_2$  exchange through the southern boundary on the offshore side ((c) in Fig. 4.6). The reduced oxygen content in the NOO2 and NOO2NO3PO4 simulations induce a more significant vertical expansion of the OMZ, with a stronger impact on the southern side and inshore. Toward the southern border, the maximum thicknesses are 3950m in these two configurations ((d) and (e) in Fig. 4.6). At around 10°S, the  $O_2$ -depleted waters are observed between 200 m and 1000 m depth, showing a vertical expansion of about 200 m. These simulations suggest a strong northward transport of oxygenated waters from the south. As the  $O_2$  concentration in the southern waters decreases, the OMZ of the ETSP might expand.

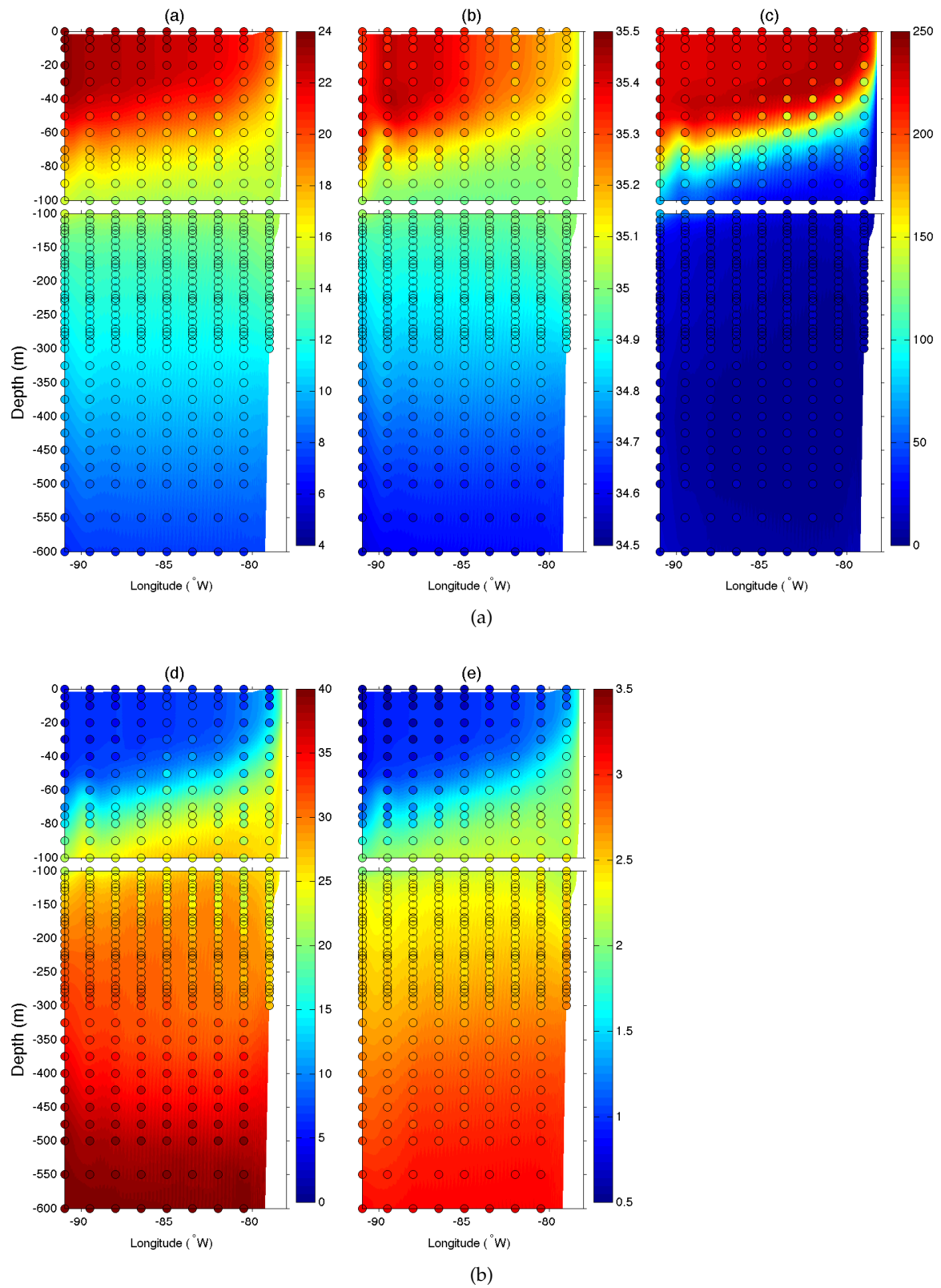


Figure 4.4: Simulated annual average (a) temperature, (b) salinity, (c) oxygen, (e) nitrate, and (d) phosphate concentrations at 10°S from surface to 600m depth, zoomed out for 0-100m depth. Colored circles represent the annual mean CARS (2009) datasets. Both model fields and data apply the same color bar in each panel.

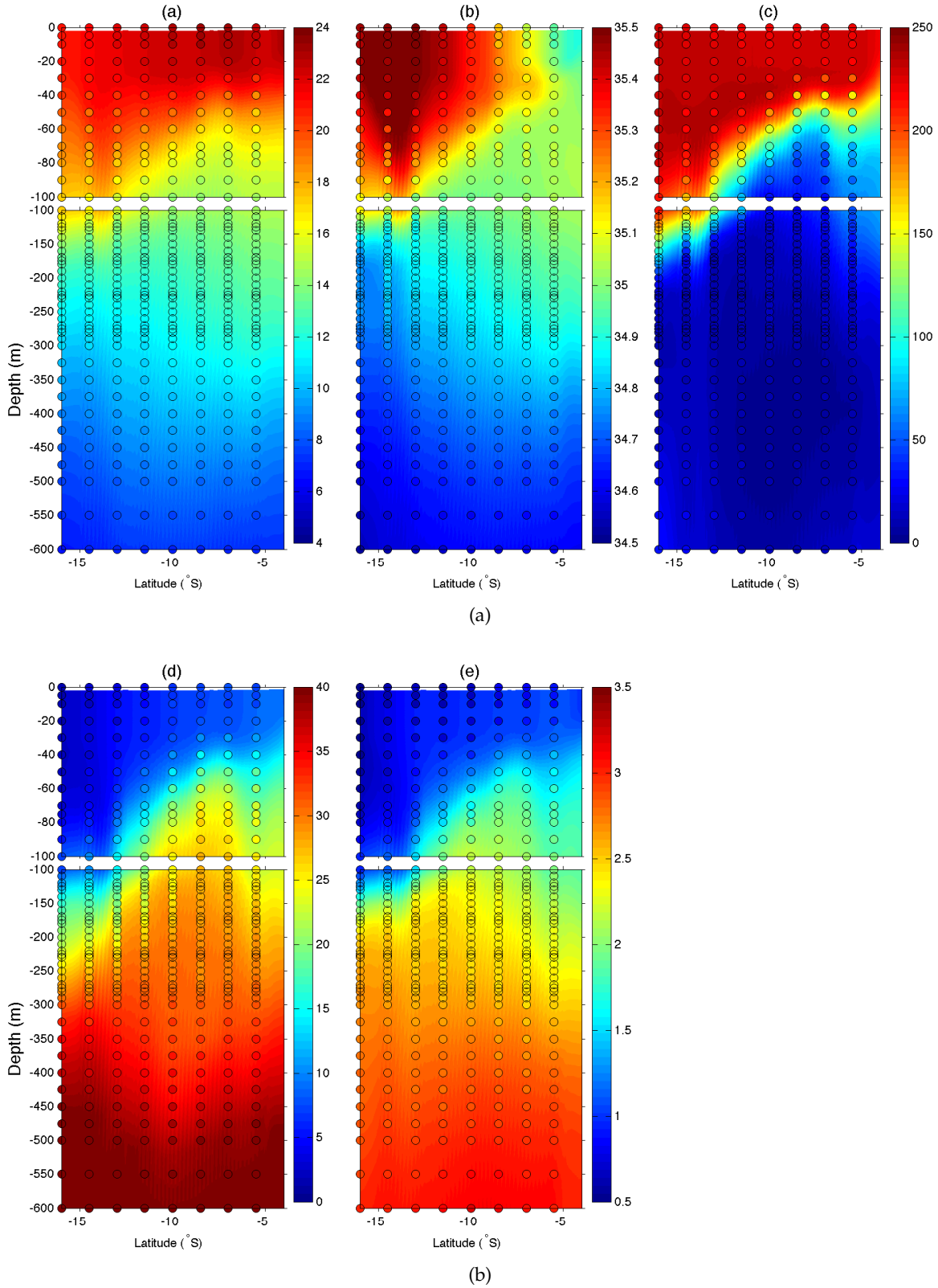


Figure 4.5: Simulated annual average (a) temperature, (b) salinity, (c) oxygen, (e) nitrate, and (d) phosphate concentrations at 85°W from surface to 600m depth, zoomed out for 0-100m depth. Colored circles represent the annual mean CARS (2009) datasets. Both model fields and data apply the same color bar in each panel.

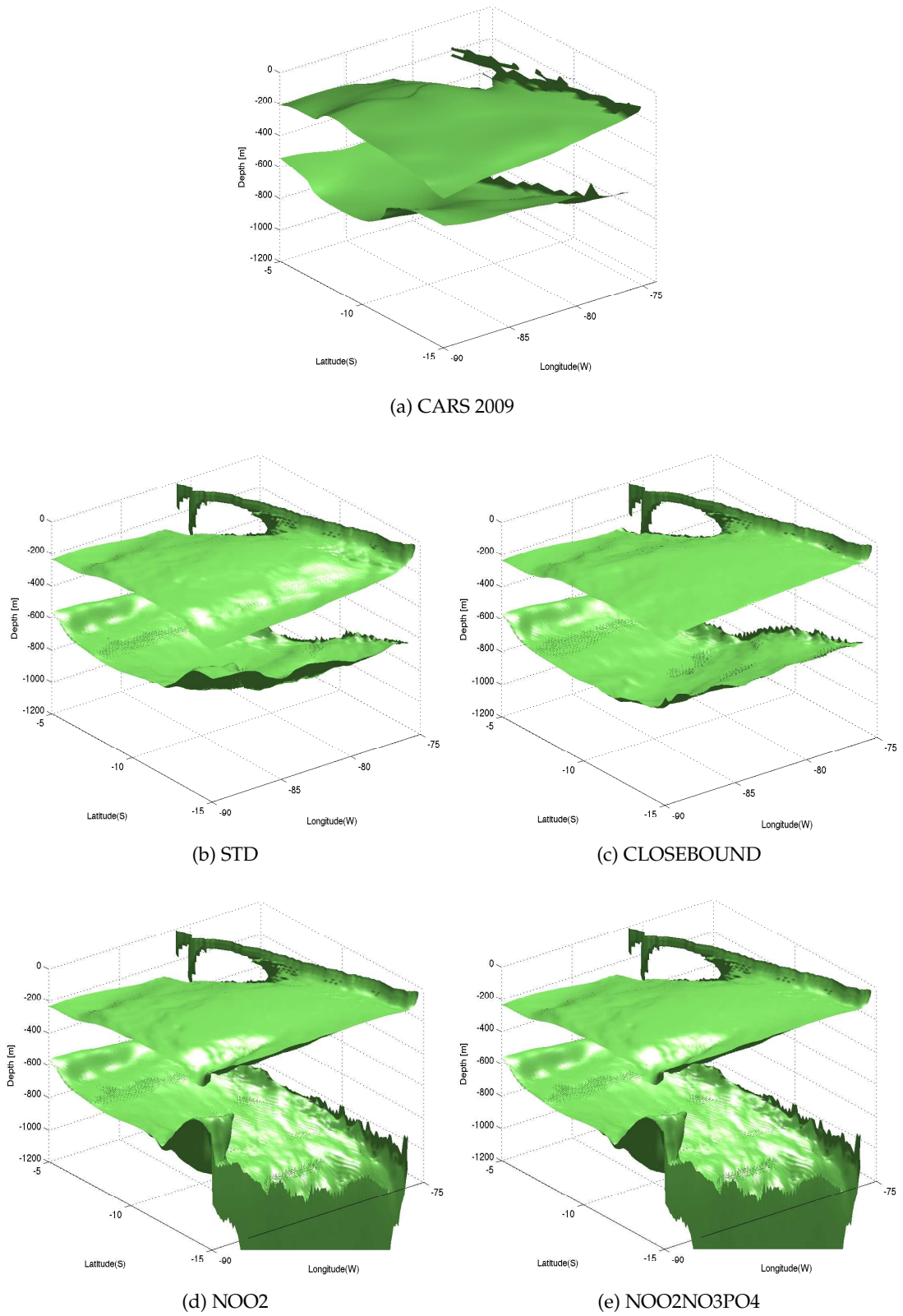


Figure 4.6: Structure evolution of the OMZ. Green surfaces represents isosurfaces with the annual average  $\text{O}_2$  concentration of  $20 \mu\text{mol kg}^{-1}$ .

### 4.3.2 O<sub>2</sub> and nutrient contents

The changes in the southern boundary conditions also modify the O<sub>2</sub> and nutrient contents of the ETSP (Fig. 4.7). We estimate both the upper 600m (to focus on the OMZ core) and the whole model domain (Fig. 4.7). The vertical profiles of the annual mean O<sub>2</sub>, NO<sub>3</sub><sup>-</sup> and PO<sub>4</sub><sup>3-</sup> concentrations are also presented in Fig. 4.8. Both contents and vertical profiles show good agreement with CARS 2009 data, indicative of good model skills in simulating the biogeochemical features of the region.

In the upper 600m, the O<sub>2</sub> and nutrient contents show generally slight changes in the idealised experiments (CLOSEBOUND, NOO2 and NOO2NO3PO4) in comparison with the STD (Figs. 4.7 and 4.8), e.g., O<sub>2</sub> content decreases by about 16%, 15% and 15% respectively in the CLOSEBOUND, NOO2 and NOO2NO3PO4 configurations; NO<sub>3</sub><sup>-</sup> content decreases by about 4.6%, 1.4% and 13% respectively; however, PO<sub>4</sub><sup>3-</sup> content decreases only in NOO2NO3PO4, stays constant in NOO2, but increases by about 5.5% in CLOSEBOUND. There are very small chlorophyll a concentration differences in the surface layer of different model configurations, except in the CLOSEBOUND configuration where there is higher chlorophyll a concentration (Fig.4.9). This can be attributed to the slightly higher NO<sub>3</sub><sup>-</sup> and PO<sub>4</sub><sup>3-</sup> concentrations at the surface of the CLOSEBOUND configuration (Fig. 4.8a).

Different from the upper layer dynamics, the net contributions of the southern boundary conditions to the oxygen and nutrient contents appear more significant in the whole water column in NOO2 and NOO2NO3PO4, but not in CLOSEBOUND ((b) in Figs. 4.7 and 4.8). For example, O<sub>2</sub> content decreases by about 36% respectively in NOO2 and NOO2NO3PO4; NO<sub>3</sub><sup>-</sup> and PO<sub>4</sub><sup>3-</sup> decrease by about 29% and 28% respectively in NOO2NO3PO4. The reduction in oxygen supplies through the southern boundary reduces the O<sub>2</sub> content, and a stronger reduction is observed in the NOO2 and NOO2NO3PO4 configurations, suggesting that a reduction of southern-origin O<sub>2</sub> might impact the OMZ of the ETSP. As for the O<sub>2</sub> dynamics in the water column, the reduction in southern-origin nutrient in NOO2NO3PO4 induce changes in the nutrient availability in the ETSP region. It is important to highlight that these changes are stronger in the deeper layers, from 600 m down to the bottom (Fig. 4.7b).

### 4.3.3 Ocean volume distributions as a function of annual mean O<sub>2</sub> and nutrient concentrations

The volume distributions as a function of O<sub>2</sub> and nutrient concentrations are used as a metric for investigating the impact of the southern boundary on the biogeochemistry of the ETSP (Fig. 4.10). At the upper 600 meter, all oxygen bins between 0 and 300 µmol l<sup>-1</sup> show weaker responses to the reduced O<sub>2</sub> supplies from the southern boundary (Fig. 4.10a), which is in agreement with Figs. 4.7a and 4.8a. The most significant change is observed in the deoxygenated wa-

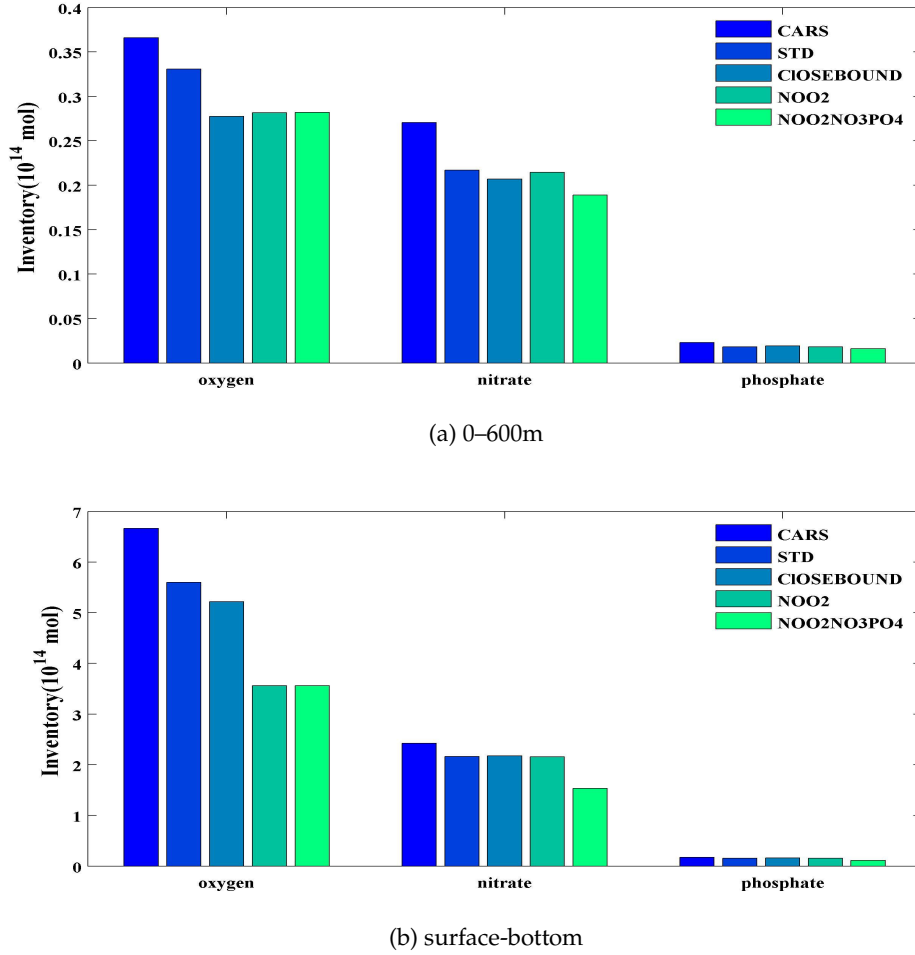
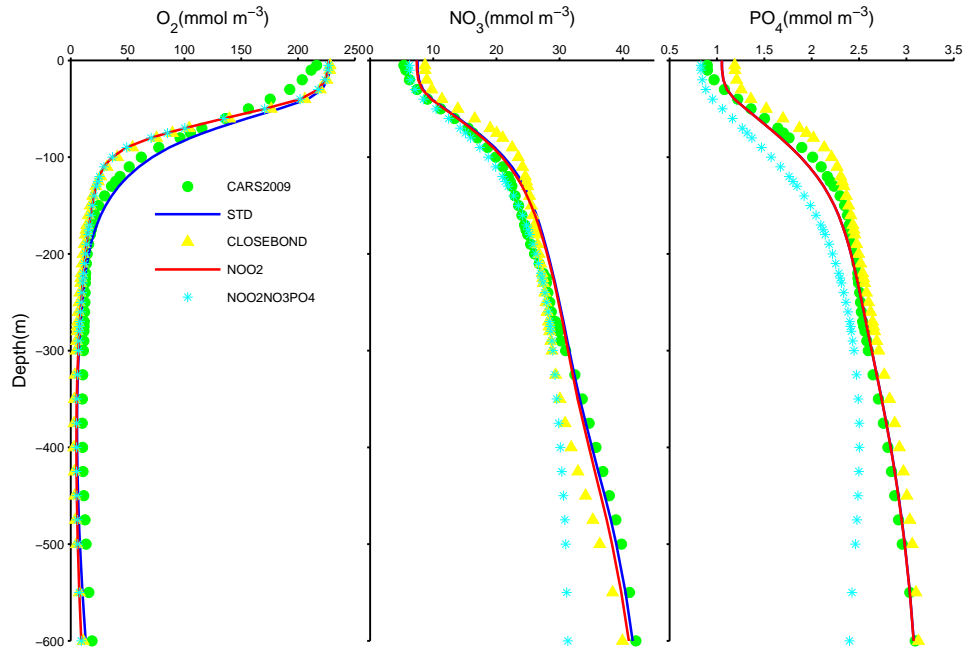


Figure 4.7: Annual mean contents of  $O_2$ ,  $NO_3$  and  $PO_4$  respectively for the model domain. The contents are estimated respectively (a) above 600m and (b) the whole water column for both CARS 2009 data and model simulations.

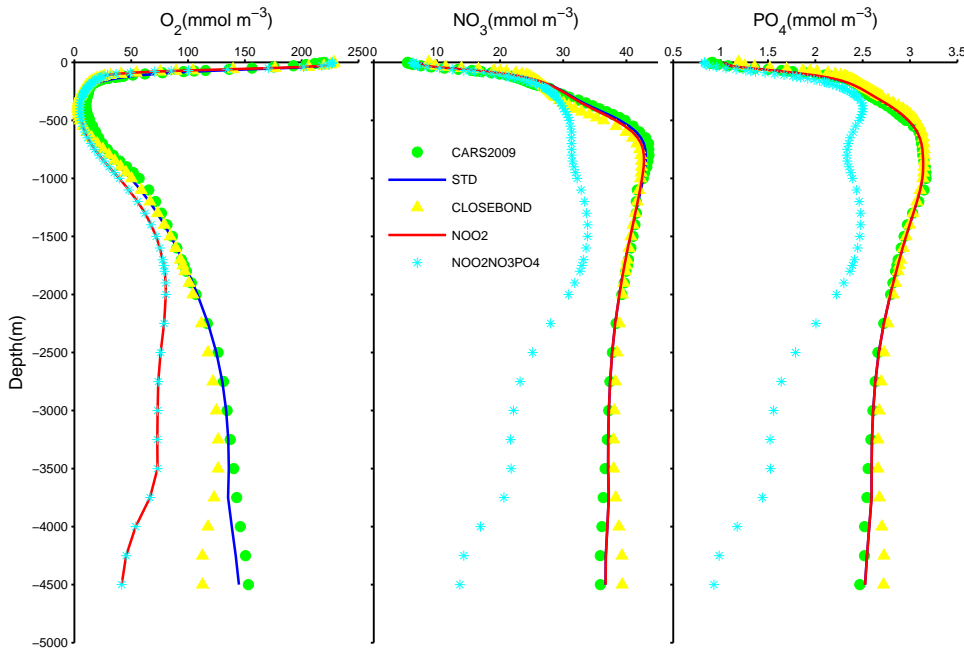
ter volume with  $O_2$  concentration between 0 and  $20 \mu\text{mol kg}^{-1}$ , as the volumes increase about 5% in the three idealised configurations (CLOSEBOUND, NOO2 and NOO2NO3PO4). For the whole model domain, the volume changes in relation to the boundary condition changes show a stronger impact, especially for the volume bins with  $O_2$  concentration between 100 and  $200 \mu\text{mol l}^{-1}$  (Fig. 4.10b). Removing the  $O_2$  supply from the southern boundary (in NOO2 and NOO2NO3PO4) leads to a decrease of as much as 44% for the oxygenated volume. These results support the suggestion that the southern boundary  $O_2$  supply mainly influences the deep ocean rather than the upper 600m.

The contributions of the southern boundary to the volume distributions of  $NO_3^-$  are variable among the simulations (Figs. 4.10c and 4.10d). In the CLOSEBOUND configuration, this distribution has been mainly changed only in the upper 600m, i.e., the bin with  $NO_3^-$  concentration between 25 and  $30 \mu\text{mol kg}^{-1}$  decreases by about 44% (Fig. 4.10c). There is no obvious impact





(a) 0–600m



(b) Surface–bottom

Figure 4.8: Simulated annual average vertical profiles for oxygen, nitrate and phosphate concentrations from surface to (a) 600m and (b) bottom respectively. Vertical profiles are average for the region of 5–15°S and 72–90°W.



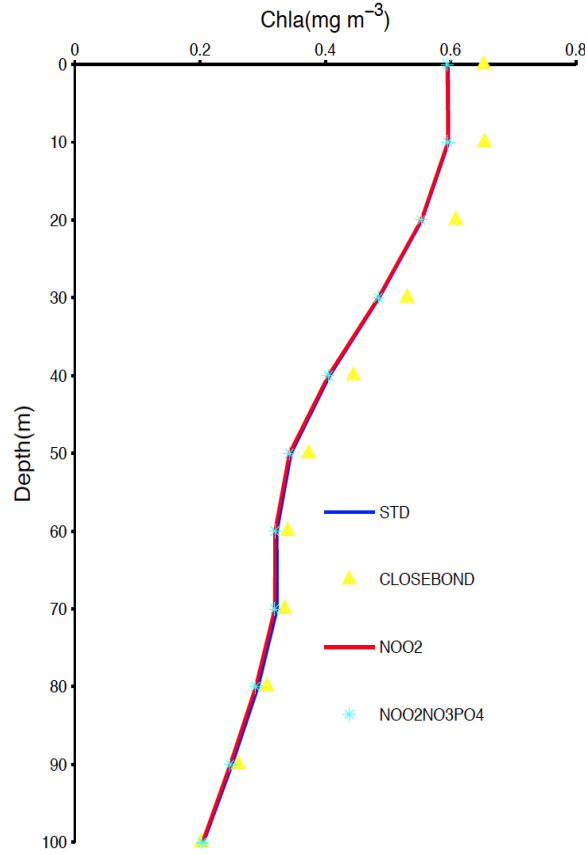


Figure 4.9: Annual mean vertical profile for chlorophyll a concentrations from the surface to 100m depth for different model configurations.

from the southern boundary conditions in the NOO2 configuration. The most significant influence is found in the NOO2NO3PO4 configuration, where the nitrate bins start to decrease from 30 and 35  $\mu\text{mol kg}^{-1}$  respectively in the upper 600m and the whole water column respectively. Generally, the effect of removing the  $\text{NO}_3^-$  supply from the southern boundary is to shift the water volumes from high concentration bins to low concentration bins.

The volume distributions of  $\text{PO}_4^{3-}$  also show variable responses to the southern boundary conditions (Figs. 4.10e and 4.10f). Compared with the STD configuration, a stronger increase in the volume is observed in the phosphate bins of 2.0-2.5  $\mu\text{mol kg}^{-1}$ , while a stronger decrease in volume distribution is observed in 2.5-3.0  $\mu\text{mol kg}^{-1}$  in both the upper 600m and the whole water column of the NOO2 and NOO2NO3PO4 configurations. Thus, in these two idealised configurations, the impact of removing  $\text{PO}_4^{3-}$  supply from the southern boundary is to reduce water volumes with  $\text{PO}_4^{3-}$  concentration more than 2.5  $\mu\text{mol kg}^{-1}$ . However, the volume increases in high  $\text{PO}_4^{3-}$  concentration bins in the CLOSEBOUND configuration, which indicates that the  $\text{PO}_4^{3-}$  content increases (Fig. 4.7). This suggests that a fully closed southern boundary can prevent  $\text{PO}_4^{3-}$  loss and subsequently increase primary production in the model domain, which is consistent with the box model result of Su et al. [2015].

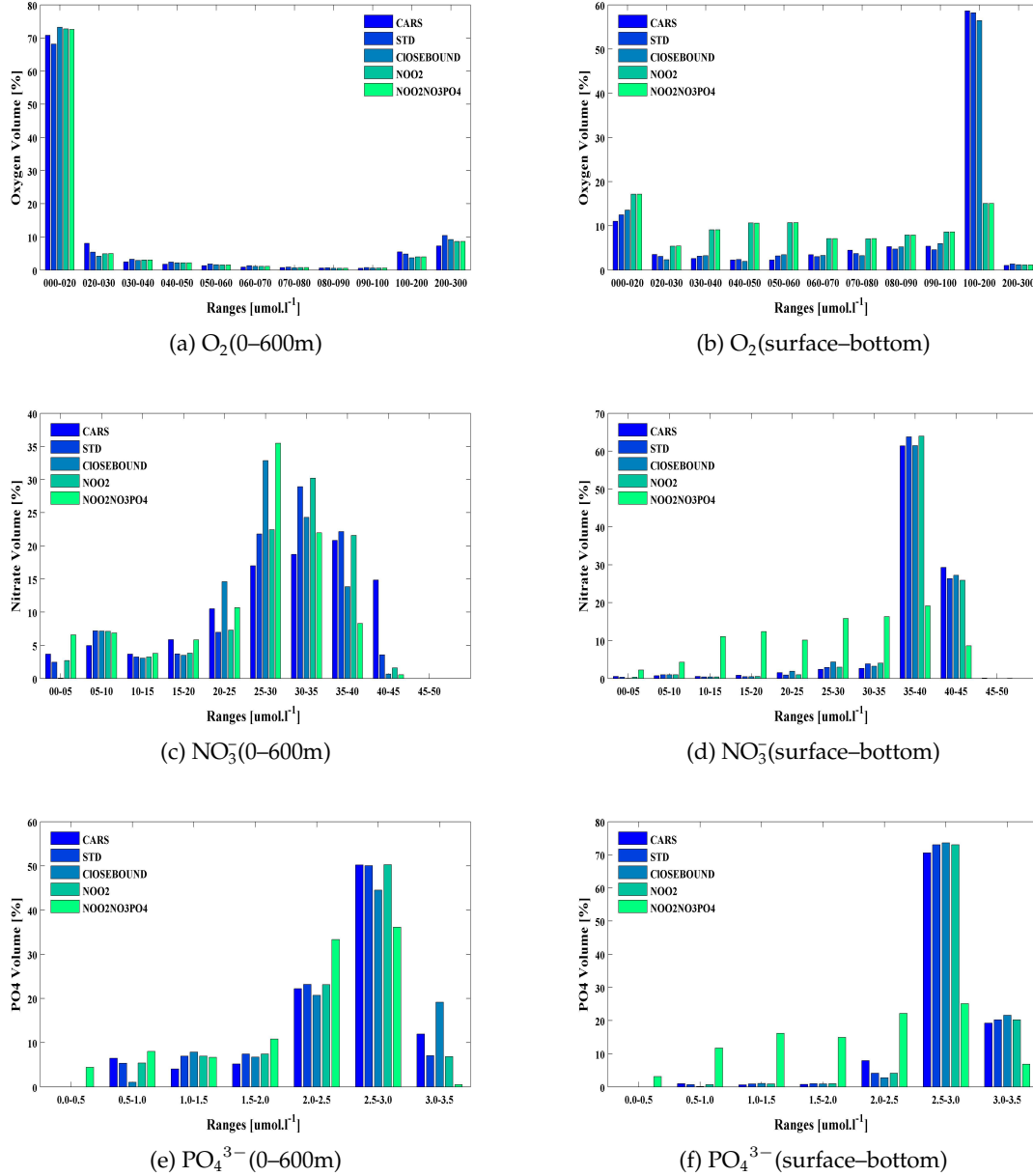


Figure 4.10: Ocean volume distributions as a function of annual mean  $O_2$ ,  $NO_3^-$  and  $PO_4^{3-}$  concentrations respectively. The contents are estimated above the 600m depth (a, c, e) and full vertical depth (b, d, f) for both CARS 2009 data and annual model means for the domain 72–90°W/5–15°S.

#### 4.3.4 Lateral fluxes

The above model assessment results show good model performance for the annual means of the ETSP, which provides the possibility to compare the fluxes through open boundaries in different model configurations. The focus will be on the lateral fluxes when analysing the  $O_2$

and nutrient concentrations of different model configurations, especially for the region below 600m.

In the upper 600m of the ocean, both  $O_2$  and nutrients are lost through the southern boundary (Table 4.1) in the STD configuration mainly due to the combined effect of the surface equatorward PCC and subsurface southward PCUC. But only little  $O_2$  is lost through the southern boundary, which can be attributed to the southward PCUC carrying OMZ water with low  $O_2$  concentration. This loss term is lacking in the CLOSEBOUND configuration, which can partially explain why  $PO_4^{3-}$  concentrations are higher in this configuration than in the STD configuration (in Figs. 4.7a and 4.10e). But this does not apply to  $NO_3^-$ , probably due to the higher  $NO_3^-$  consumption via denitrification resulting from higher export production (Fig. 4.9). In the NOO2 and NOO2NO3PO4 configurations, there is much more  $O_2$  loss through the southern boundary, but there are only slight  $O_2$  variations (Fig. 4.8a), which can be attributed to both less loss through the western boundary and more input from the atmosphere (Table 4.1). In the NOO2NO3PO4 configuration,  $NO_3^-$  and  $PO_4^{3-}$  concentrations are lower than those in the STD configuration (Fig. 4.8a), resulting in less  $NO_3^-$  and  $PO_4^{3-}$  losses from the southern boundary.

The entire model domain gains  $O_2$  through the southern boundary (Table 4.1).  $O_2$  is lost through the southern boundary in the upper 600m and the PCDCC is the main equatorward current below 600m. Therefore, this gain can only be due to the equatorward PCDCC carrying  $O_2$  rich water into the ETSP. In the CLOSEBOUND configuration in which all the connection with the southern subtropical Pacific is closed, less  $NO_3^-$  and  $PO_4^{3-}$  are lost through the southern boundary (Table 4.1). Since there is more loss through the northern boundary and less input of  $NO_3^-$  and  $PO_4^{3-}$  from the western boundary (Table 4.1), the slightly higher contents of these two nutrients in the whole domain can only be attributed to the southern boundary (Fig. 4.7b). The higher concentrations in the deep ocean may be due to the nutrients imported via nutrient-rich southward PDW (Fig. 4.8b). In the NOO2 and NOO2NO3PO4 configurations, there are more  $O_2$  losses from the southern boundaries (Table 4.1), mainly resulting in low  $O_2$  concentrations in the model domain (Fig. 4.8b). The lower  $NO_3^-$  and  $PO_4^{3-}$  concentrations below the subsurface in the model domain (Fig. 4.8b) can be attributed to the higher  $NO_3^-$  and  $PO_4^{3-}$  losses through the southern boundary (Table 4.1).

## 4.4 Conclusions

We used a coupled physical-biogeochemical model to investigate the impact of the southern boundary on the  $O_2$  and nutrient variabilities in the OMZ off Peru and the ETSP. Model assessment indicates good performance of the model in simulating the main physical and biogeochemical features of the region. The influence of the southern boundary on the OMZ is investigated in terms of  $O_2$ ,  $NO_3^-$  and  $PO_4^{3-}$  by analysing model configurations with different southern boundary conditions.

STD				CLOSEBOUND			NOO2			NOO2NO3PO4		
0-600m												
	O <sub>2</sub>	NO <sub>3</sub> <sup>-</sup>	PO <sub>4</sub> <sup>3-</sup>	O <sub>2</sub>	NO <sub>3</sub> <sup>-</sup>	PO <sub>4</sub> <sup>3-</sup>	O <sub>2</sub>	NO <sub>3</sub> <sup>-</sup>	PO <sub>4</sub> <sup>3-</sup>	O <sub>2</sub>	NO <sub>3</sub> <sup>-</sup>	PO <sub>4</sub> <sup>3-</sup>
south	-0.721	-30.2	-2.54	0	0	0	-51.9	-29.7	-2.54	-52.5	-21.7	-1.92
north	136	18.3	1.68	124	11.1	1.11	136	18.3	1.68	136	18.4	1.68
west	-247	32.3	1.96	-230	15.2	0.396	-205	32.4	1.96	-205	34.2	2.31
Atm	2.87			7.59			46.8			46.6		
sum	-109	20.5	1.10	-98.5	26.3	1.50	-74.1	21.0	1.11	-74.7	30.9	2.07
	surface-bottom											
	O <sub>2</sub>	NO <sub>3</sub> <sup>-</sup>	PO <sub>4</sub> <sup>3-</sup>	O <sub>2</sub>	NO <sub>3</sub> <sup>-</sup>	PO <sub>4</sub> <sup>3-</sup>	O <sub>2</sub>	NO <sub>3</sub> <sup>-</sup>	PO <sub>4</sub> <sup>3-</sup>	O <sub>2</sub>	NO <sub>3</sub> <sup>-</sup>	PO <sub>4</sub> <sup>3-</sup>
south	45.4	-26.8	-2.38	0	0	0	-78.4	-26.4	-2.38	-78.9	-33.8	-2.79
north	46.5	-7.07	-0.120	48.4	-13.7	-0.666	49.5	-7.06	-0.120	49.5	-6.21	-0.0598
west	-159	52.9	3.49	-102	39.0	2.09	-99.7	53.1	3.49	-99.5	58.5	4.07
Atm	2.87			7.59			46.8			46.6		
sum	-64.6	19.1	0.996	-45.8	25.3	1.43	-81.7	19.7	0.996	-82.4	18.4	1.22

Table 4.1: Oxygen and nutrient fluxes through boundaries via horizontal advections and air-sea gas exchange ( $10^{11} \text{ mol yr}^{-1}$ ). “Atm” is the oxygen flux through sea surface. Positive values indicate gains into the model domain, while negative values losses.

The southern boundary conditions appear to have significant influence on the O<sub>2</sub> distribution in the ETSP. Even though the lack of O<sub>2</sub> input from the southern subtropical Pacific can enlarge the OMZ, the impacts on critical O<sub>2</sub> distribution and concentration changes are mainly in the ocean deeper than 600m (Figs. 4.10b and 4.8b). The O<sub>2</sub> input from atmosphere plays an important role in compensating for lack of O<sub>2</sub> input from the southern boundary at the upper layer (Table 4.1).

The ETSP is a gaining both NO<sub>3</sub><sup>-</sup> and PO<sub>4</sub><sup>3-</sup> from the adjacent ocean. But both NO<sub>3</sub><sup>-</sup> and PO<sub>4</sub><sup>3-</sup> are lost to the adjacent subtropical ocean through the southern boundary. Compared with the CLOSEBOUND configuration, our model domain loses NO<sub>3</sub><sup>-</sup> and PO<sub>4</sub><sup>3-</sup> in the realistic STD configuration when the southern boundary is open. This corroborates each other with the box model of Su et al. [2015]. In the upper layer, removing NO<sub>3</sub><sup>-</sup> and PO<sub>4</sub><sup>3-</sup> supplies from the southern boundary can also reduce the net loss in all idealised model configurations.

By changing the boundary conditions, it is evident that the northward water originating from

south into the OMZ off Peru can induce changes in the OMZ thickness and  $O_2$  and nutrient distributions. The stronger changes of oxygen are mainly observed in the deeper layers, suggesting a pathway of deep northward intrusion of well-oxygenated southern waters.

$O_2$  concentrations are critical for fish and marine animals. The ETSP is thought to contribute more than 10% of the global fish catches [Chavez et al., 2008], thus the  $O_2$  concentrations are very important for maintaining the high fish production. Expansion of the OMZs will narrow down the habitable depth range of fishes and, together with overfishing, may threaten the sustainability of pelagic fisheries and marine ecosystems [Stramma et al., 2012]. The Southern Ocean is thought to be sensitive to global warming due to reductions in deep convection, resulting in a significant decrease in  $O_2$  concentrations [Keeling and Garcia, 2002, and references therein]. The Southern Ocean is an important source of water masses in the ETSP. Thus, this study can give some insights into the impacts of Southern Ocean variation on the oxygen cycle of the ETSP.

## 4.5 Appendix A

A 30-year simulation is conducted to attain a nearly repeating annual cycle. Figures 4.11 and 4.12 depicts the time variations of both physical and biogeochemical variables. For each of these variables, after a spin-up of 25 years, the model exhibits no temporal drift, especially the volume-averaged kinetic energy (Fig. 4.11) and nitrate concentrations (Fig. 4.12). Afterwards, they stay stable.

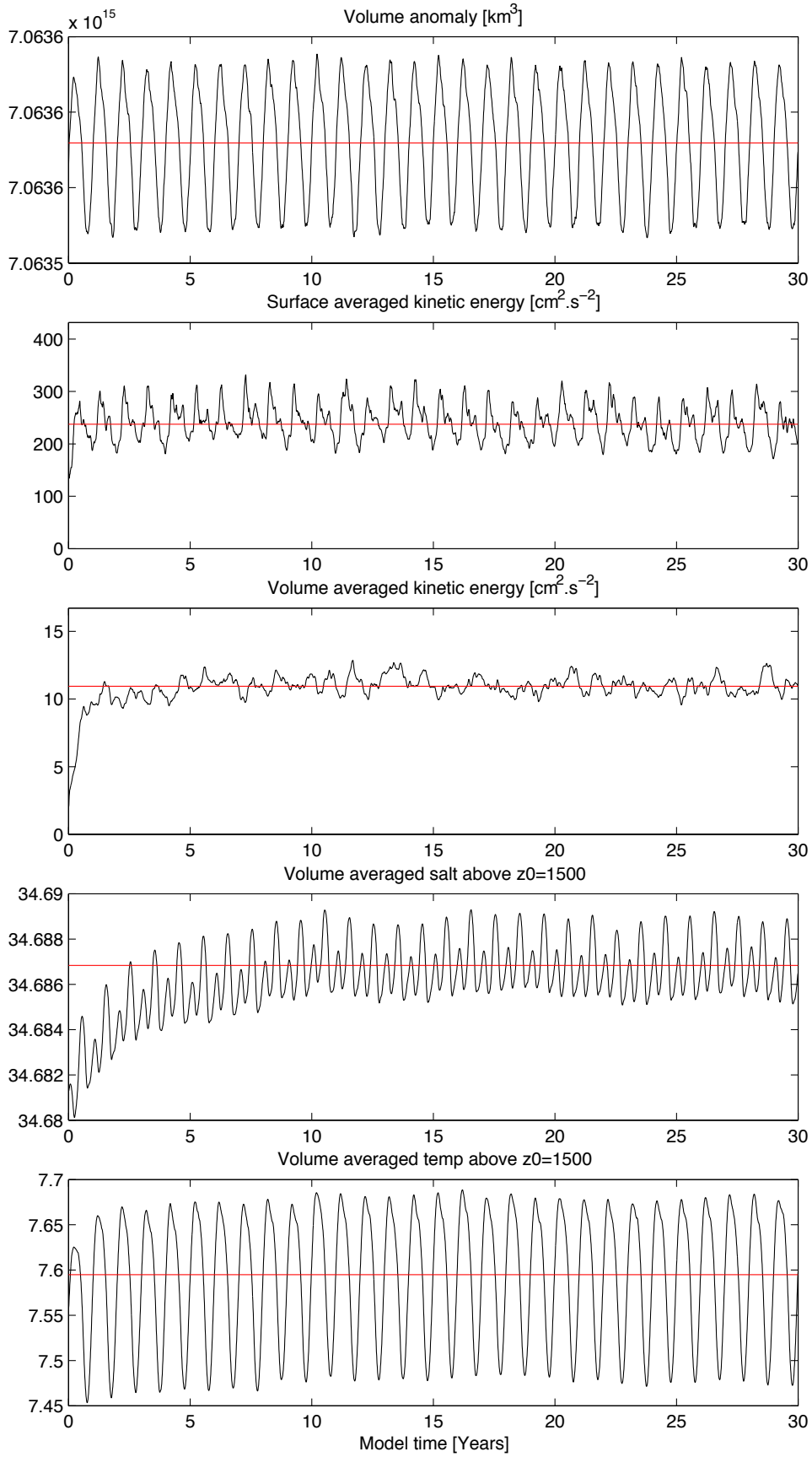


Figure 4.11: Time evolution of simulated physical variables.

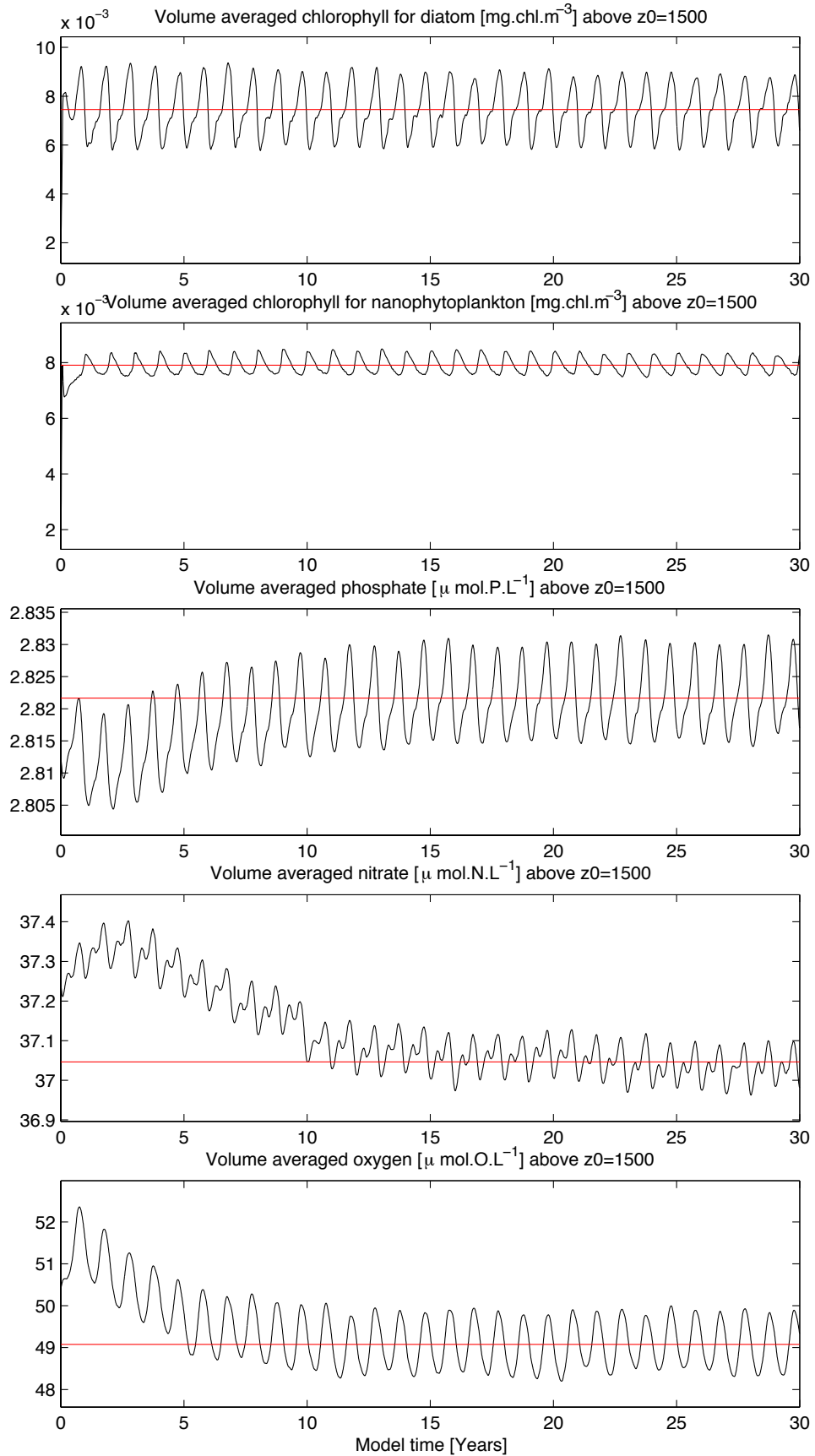


Figure 4.12: Time evolution of simulated biogeochemical variables.





## Chapter 5

# Summary and outlook

The nitrogen cycle in the Oxygen Minimum Zone (OMZ) of the eastern tropical South Pacific (ETSP) has been simulated with both 2-D box and 3-D coupled physical-biogeochemical models. In the 2-D box model, reduced remineralization by denitrification under anoxic conditions is found to be responsible for preventing nitrogen depletion in the OMZ of the ETSP [Su et al., 2015]. After including the atmospheric deposition and benthic remineralisation, the local responses indicate dominant stabilizing feedbacks in the ETSP, tending to keep a balanced nitrogen inventory. All of these results indicate possible mechanisms that are responsible for keeping a stabilized nitrogen inventory in the OMZ of the ETSP. Under the current notion of nitrogen fixation being favoured in nitrogen-deficit waters, the model domain of the ETSP is a  $\text{NO}_3^-$  source. After including benthic denitrification, the ETSP might become a  $\text{NO}_3^-$  sink. We also find in the box model that  $\text{O}_2$  supply into the model domain from the southern boundary is very important in retrieving the realistic  $\text{O}_2$  concentration in the deep ocean.

Based on the important role of the southern boundary on the oxygen and nutrient contents of the ETSP found in the box model [Su et al., 2015], a model configuration of the same region is built with the high-resolution hydrodynamic model ROMS (*Regional Ocean Modelling System*) [Shchepetkin and McWilliams, 2005], coupled with the biogeochemical model PISCES (*Pelagic Interaction Scheme for Carbon and Ecosystem Studies*) [Aumont and Bopp, 2006]. A validation against in situ observations shows a realistic simulation of the horizontal and vertical oxygen and nutrient distributions by the standard configuration. Comparisons among different model configurations indicate that the southern boundary has significant influence on  $\text{O}_2$  and nutrient contents and distributions of this region. Lateral fluxes estimations indicate that our model domain is gaining  $\text{O}_2$  and losing  $\text{NO}_3^-$  and  $\text{PO}_4^{3-}$  through the southern boundary. The southern boundary is found to have more significant influence on  $\text{O}_2$  concentration of the deep layer than that of the upper layer, suggesting a pathway of deep northward intrusion of well-oxygenated southern waters. The Southern Ocean is thought to be sensitive to global warming due to reductions in deep convection, resulting in significant decrease in  $\text{O}_2$  con-

centrations. This study can give insights into the impacts of Southern Ocean variations due to climate change on the  $O_2$  changes of the ETSP.

In the high-resolution coupled physical-biogeochemical model, a stabilised nitrogen inventory is observed in the OMZ of the ETSP even though remineralisation rates are identical under oxic and anoxic conditions. Dissolved organic matter (DOM) is found to be exported out of model domain. This could explain the stabilised nitrogen inventory, because DOM cycling is thought to be capable of promoting the spacial decoupling of nitrogen fixation and denitrification, and allowing for negative feedbacks stabilising the nitrogen inventory [Landolfi et al., 2013]. A future research topic could be to further investigate the mechanisms capable of preventing the essential complete drawdown of  $NO_3^-$  in the high-resolution coupled physical-biogeochemical model.

The interannual variations in the Pacific (i.e., El Niño-Southern Oscillation, ENSO) have an important role on the climate and marine primary production of the ETSP [Wang and Fiedler, 2006]. The circulation and water-mass connections are also found to change during the ENSO period [Montes et al., 2011]. How dose the nitrogen inventory in this region vary corresponding to the ENSO-induced variations? What is the nitrogen stabilising mechanism in the ENSO period? These questions can potentially also be answered using the high-resolution coupled physical-biogeochemical model applied in this dissertation.

To have better understanding of the full nitrogen cycle in the ETSP, benthic remineralization processes can be included using Bohlen et al. [2012]’s transfer function and the relation for phosphate regeneration following Wallmann [2010] in the high-resolution model. The interactions among nitrogen fixation, water-column and benthic denitrification, and the question whether the ETSP is a fixed-nitrogen source or sink can be further investigated.

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# Declaration

Bei Su, declares that the design and content of this thesis, apart from my supervisors' guidance, are my own work except as stated below:

Computation setting in the computing-station of the Kiel University, scientific suggestions from all my coauthors for the published and submitted manuscripts.

The thesis has not been submitted either partially or wholly to apply for doctoral degree to any other examining body. And it has been prepared following the Rules of Good Scientific Practice of the German Research Foundation.

Kiel, April 27, 2015

Bei Su

